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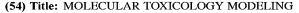
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(57) Abstract: The present invention is based on the elucidation of the global changes in gene expression and the identification of toxicity markers in tissues or cells exposed to a known renal toxin. The genes may be used as toxicity markers in drug screening and toxicity assays. The invention includes a database of genes characterized by toxin-induced differential expression that is designed for use with microarrays and other solid-phase probes.





MOLECULAR TOXICOLOGY MODELING

INVENTORS: Donna MENDRICK, Mark PORTER, Kory JOHNSON, Brandon HIGGS, Arthur CASTLE, and Michael ELASHOFF

RELATED APPLICATIONS

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This application claims priority to U.S. Provisional Applications 60/292,335; 60/297,523; 60/298,925; 60/303,810; 60/303,807; 60/303,808; 60/315,047; 60/324,928; 60/330,867; 60/330,462; 60/331,805; 60/336,144; 60/340,873; 60/357,843; 60/357,842; 60/357,844; 60/364,134; 60/370,206; 60/370,247; 60/370,144; 60/371,679; and 60/372,794, all of which are herein incorporated by reference in their entirety. This application is also related to U.S. Application Nos. 09/917,800 and 10/060,087, both of which are also herein incorporated by reference in their entirety.

SEQUENCE LISTING SUBMISSION ON COMPACT DISC

The Sequence Listing submitted concurrently herewith on compact disc is herein incorporated by reference in its entirety. Four copies of the Sequence Listing, one on each of four compact discs are provided. Copy 1, Copy 2 and Copy 3 are identical. Copies 1, 2, and 3 are also identical to the CRF. Each electronic copy of the Sequence Listing was created on May 22, 2002 with a file size of 3088 KB. The file names are as follows: Copy 1- gl5089wo.txt; Copy 2- gl5089wo.txt; CRF- gl5089wo.txt.

BACKGROUND OF THE INVENTION

The need for methods of assessing the toxic impact of a compound, pharmaceutical agent or environmental pollutant on a cell or living organism has led to the development of procedures which utilize living organisms as biological monitors. The simplest and most convenient of these systems utilize unicellular microorganisms such as yeast and bacteria, since they are the most easily maintained and manipulated. In addition, unicellular screening systems often use easily detectable changes in phenotype to monitor the effect of test compounds on the cell. Unicellular organisms, however, are inadequate models for estimating the potential effects of many compounds on complex multicellular animals, as they do not have the ability to carry out biotransformations.

The biotransformation of chemical compounds by multicellular organisms is a significant factor in determining the overall toxicity of agents to which they are exposed. Accordingly, multicellular screening systems may be preferred or required to detect the

toxic effects of compounds. The use of multicellular organisms as toxicology screening tools has been significantly hampered, however, by the lack of convenient screening mechanisms or endpoints, such as those available in yeast or bacterial systems. Additionally, previous attempts to produce toxicology prediction systems have failed to provide the necessary modeling data and statistical information to accurately predict toxic responses (e.g., WO 00/12760, WO 00/47761, WO 00/63435, WO 01/32928, and WO 01/38579).

SUMMARY OF THE INVENTION

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The present invention is based on the elucidation of the global changes in gene expression in tissues or cells exposed to known toxins, in particular renal toxins, as compared to unexposed tissues or cells as well as the identification of individual genes that are differentially expressed upon toxin exposure.

In various aspects, the invention includes methods of predicting at least one toxic effect of a compound, predicting the progression of a toxic effect of a compound, and predicting the renal toxicity of a compound. The invention also includes methods of identifying agents that modulate the onset or progression of a toxic response. Also provided are methods of predicting the cellular pathways that a compound modulates in a cell. The invention also includes methods of identifying agents that modulate protein activities.

In a further aspect, the invention includes probes comprising sequences that specifically hybridize to genes in Tables 1-5. Also included are solid supports comprising at least two of the previously mentioned probes. The invention also includes a computer system that has a database containing information identifying the expression level in a tissue or cell sample exposed to a renal toxin of a set of genes comprising at least two genes in Tables 1-5.

DETAILED DESCRIPTION

Many biological functions are accomplished by altering the expression of various genes through transcriptional (e.g. through control of initiation, provision of RNA precursors, RNA processing, etc.) and/or translational control. For example, fundamental biological processes such as cell cycle, cell differentiation and cell death, are often characterized by the variations in the expression levels of groups of genes.

Changes in gene expression are also associated with the effects of various chemicals, drugs, toxins, pharmaceutical agents and pollutants on an organism or cell. For example, the lack of sufficient expression of functional tumor suppressor genes and/or the over expression of oncogene/protooncogenes after exposure to an agent could lead to tumorgenesis or hyperplastic growth of cells (Marshall (1991), *Cell* 64: 313-326; Weinberg (1991), *Science* 254: 1138-1146). Thus, changes in the expression levels of particular genes (*e.g.* oncogenes or tumor suppressors) may serve as signposts for the presence and progression of toxicity or other cellular responses to exposure to a particular compound.

Monitoring changes in gene expression may also provide certain advantages during drug screening and development. Often drugs are screened for the ability to interact with a major target without regard to other effects the drugs have on cells. These cellular effects may cause toxicity in the whole animal, which prevents the development and clinical use of the potential drug.

The present inventors have examined tissue from animals exposed to known renal toxins which induce detrimental kidney effects, to identify global changes in gene expression induced by these compounds. These global changes in gene expression, which can be detected by the production of expression profiles (an expression level of one or more genes), provide useful toxicity markers that can be used to monitor toxicity and/or toxicity progression by a test compound. Some of these markers may also be used to monitor or detect various disease or physiological states, disease progression, drug efficacy, and drug metabolism.

Identification of Toxicity Markers

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To evaluate and identify gene expression changes that are predictive of toxicity, studies using selected compounds with well characterized toxicity have been conducted by the present inventors to catalogue altered gene expression during exposure *in vivo* and *in vitro*. In the present study, cephaloridine, cisplatin, puromycin aminonucleoside (PAN), bromoethylamine hydrobromide (BEA), gentamicin, ifosfamide, cyclophosphamide, carboplatin, AY-25329, indomethacin, acyclovir, citrinin, mercuric chloride, diflunisal, cidofovir, pamidronate, lithium, hydralazine, colchicine, sulfadiazine, and adriamycin were selected as known renal toxins.

Cephaloridine is an amphoteric, semi-synthetic, broad-spectrum cephalosporin derived from cephalosporin C. Cephalosporins are β-lactam-containing antibiotics which prevent bacterial growth by inhibiting polymerization of the peptidoglycan bacterial cell wall. The linear glycan chains (composed of N-acetylglucosime and N-acetylmuramic acid) are cross-linked to each other by the coupling of short chains of several amino acids, the coupling resulting from the action of a transpeptidase. It is believed that cephalosporins act by blocking the activity of the transpeptidase (Goodman & Gilman's The Pharmalogical Basis of Therapeutics 9th ed., J.G. Hardman *et al.* Eds., McGraw Hill, New York, 1996, pp. 1074-1075, 1089-1095).

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Cephaloridine is administered intramuscularly and is used to treat infections of the respiratory tract, gastrointestinal tract and urinary tract, as well as infections of soft tissue, bones and joints. Noted adverse effects include hypersensitivity reactions (such as anaphylactic shock, urticaria and bronchospasm), gastrointestinal disturbances, candidiasis, and cardiovascular and blood toxicity, in particular, toxicity to the hematopoietic system (cells responsible for the formation of red and white blood cells and platelets).

Although cephaloridine may be nephrotoxic at high dosages, it is not as harmful to the kidneys as are the aminoglycosides and polymixins. High dosages of cephaloridine may cause acute renal tubular necrosis (Cecil Textbook of Medicine, 20th ed., part XII, p. 586, J. C. Bennett and F. Plum Eds., W. B. Saunders Co., Philadelphia, 1996) or drug-induced interstitial nephritis, which is accompanied by elevated IgE levels, fever, arthralgia and maculopapular rash. Renal biopsopy demonstrates edema and interstitial inflammatory lesions, mainly with lymphocytes, monocytes, eosinophils and plasma cells. Vasculitis of small vessels may develop, leading to necrotising glomerulonephritis (G. Koren, "The nephrotoxic potential of drugs and chemicals. Pharmacological basis and clinical relevance.," *Med Toxicol Adverse Drug Exp* 4(1):59-72, 1989).

Cephaloridine has also been shown to reduce mitochondrial respiration and uptake of anionic succinate and carrier-mediated anionic substrate transport (Tune *et al.* (1990), *J Pharmacol Exp Ther* 252: 65-69). In a study of oxidative stress and damage to kidney tissue, cephaloridine depleted reduced glutathione (GSH) and produced oxidized glutathione (GSSG) in the renal cortex. This drug also inhibited glutathione reductase

and produced malondialdehyde and conjugated dienes (Tune et al. (1989), Biochem Pharmacol 38: 795-802). Because cephaloridine is actively transported into the proximal renal tubule, but slowly transported across the lumenal membrane into the tubular fluid, high concentrations can accumulate and cause necrosis. Necrosis can be prevented by administering inhibitors of organic anion transport, although such treatment may be counterproductive, as cephaloridine is passed in and out of the kidney by the renal organic anion transport system (Tune et al. (1980), J Pharmacol Exp Ther 215: 186-190).

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Cisplatin (Pt (NH₃)₂(Cl)₂), a broad-spectrum anti-tumor agent, is commonly used to treat tumors of the testicles, ovaries, bladder, skin, head and neck, and lungs (<u>PDR 47th ed.</u>, pp. 754-757, Medical Economics Co., Inc., Montvale, NJ, 1993; <u>Goodman & Gilman's The Pharmalogical Basis of Therapeutics 9th ed.</u>, pp. 1269-1271, J.G. Hardman *et al.* Eds., McGraw Hill, New York, 1996). Cisplatin diffuses into cells and functions mainly by alkylating the N⁷ of guanine, a highly reactive site, causing interstrand and intrastrand crosslinks in the DNA that are lethal to cells. The drug is not sensitive to the cell cycle, although its effects are most pronounced in S phase.

Because the drug is cleared from the body mainly by the kidneys, the most frequent adverse effect of cisplatin usage is nephrotoxicity, the severity of which increases with increasing dosage and treatment terms. Other adverse effects include renal tubule damage, myelosuppression (reduced numbers of circulating platelets, leukocytes and erythrocytes), nausea and vomiting, ototoxicity, serum electrolyte disturbances (decreased concentrations of magnesium, calcium, sodium, potassium and phosphate, probably resulting from renal tubule damage), increased serum concentrations of urea and creatinine, and peripheral neuropathies.

In one study on rats (Nonclercq et al. (1989), Exp Mol Pathol 51: 123-140) administration of cisplatin or carboplatin induced renal injury, carboplatin causing less damage than cisplatin. The most prominent injury was to the straight portion of proximal renal tubule.

In another rat study (Goldstein *et al.* (1981), *Toxicol Appl Pharmacol* 60: 163-175) animals injected with cisplatin displayed decreased food intake as drug dosage increased. On day 2, the high-dose groups (10-15 mg/kg) exhibited a six or seven-fold elevation in BUN. On day 4, BUN elevation was noted in the 5mg/kg group. An

increase in urine volume was observed beginning on days 3-4, along with decreased urine osmolality in the low-dose groups (2.5 or 5 mg/kg). Another experiment on rats (Agarwal *et al.* (1995), *Kidney Int* 48: 1298-1307) showed that cisplatin treatment produced elevations in serum creatinine levels, which began on day 3 and progressed for the duration of the study.

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PAN (C₂₂H₂₉N₇O₅), an antibiotic produced by *Streptomyces alboniger*, inhibits protein synthesis and is commonly used experimentally on rats to mimic human minimal change disease. One study showed that PAN-injected rats demonstrated an increase in levels of serum non-esterified fatty acids, while the serum albumin concentration was negatively affected (Sasaki *et al.* (1999), *Adv Exp Med Biol* 467: 341-346).

In another rat study, an adenosine deaminase inhibitor prevented PAN nephrotoxicity, indicating that PAN toxicity is linked to adenosine metabolism (Nosaka et al. (1997), Free Radic Biol Med 22: 597-605). Another group showed that PAN, when administered to rats, led to proteinuria, a condition associated with abnormal amounts of protein in the urine, and renal damage, e.g. blebbing of glomerular epithelial cells, focal separation of cells from the glomerular basement membrane, and fusion of podocytes (Olson et al. (1981), Lab Invest 44: 271-279). In another study on rats, administration of PAN induced glomerular epithelial cell apoptosis in a dose- and time-dependent manner (Sanwal et al. (2001), Exp Mol Pathol 70: 54-64).

One study with PAN-injected rats (Koukouritaki et al. (1998), J Investig Med 46: 284-289) examined the changes in the expression of the proteins paxillin, focal adhesion kinase, and Rho, all of which regulate cell adhesion to the extracellular matrix. Paxillin levels increased steadily, peaked at day 9 after PAN injection, and then remained elevated even after proteinuria resolved. There was no observed change in expression of either focal adhesion kinase or Rho.

BEA, (C₂H₆BrN.HBr), is commonly used experimentally on rats to induce papillary necrosis and renal cortex damage, which is similar to human analgesic nephropathy. BEA-induced papillary necrosis in rats eventually leads to the onset of focal glomerular sclerosis and nephrotic proteinuria (Garber *et al.* (1999), *Am J Kidney Dis* 33: 1033-1039). Even at low doses (50 mg/kg), BEA can induce an apex limited renal papillary necrosis (Bach *et al.* (1983), *Toxicol Appl Pharmacol* 69: 333-344). In male Wistar rats, BEA administered at 100 mg/kg was shown to cause renal papillary

necrosis within 24 hours (Bach et al. (1991), Food Chem Toxicol 29: 211-219). Additionally, Bach et al. showed that there was an increase in urinary triglycerides, and lipid deposits were seen by Oil Red O lipid staining in the cells of the collecting ducts and hyperplastic urothelia adjacent to the necrosed region.

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It has also been shown that succinate and citrate concentrations are significantly lower in the urine of BEA-treated rats (Holmes *et al.* (1995), *Arch Toxicol* 70: 89-95). Moreover, BEA treatment induced glutaric and adipic aciduria, which is symptomatic of an enzyme deficiency in the acyl CoA dehydrogenases. The same study examined urinary taurine levels in desert mice, and in BEA-treated desert mice there was an increase in the urinary taurine level which is indicative of liver toxicity.

Another study on BEA-treated rats showed that there was an increase in the concentrations of creatine in the renal papilla and glutaric acid in the liver, renal cortex, and renal medulla as soon as 6 hours post-treatment (Garrod *et al.* (2001), *Magn Reson Med* 45: 781-790).

Discovered and purified in the early 1960's, gentamicin is a broad-spectrum aminoglycoside antibiotic that is cidal to aerobic gram-negative bacteria and commonly used to treat infections, e.g., those of the urinary tract, lungs and meninges. As is typical for an aminoglycoside, the compound is made of two amino sugar rings linked to a central aminocyclitol ring by glycosidic bonds. Aminoglycosides are absorbed poorly with oral administration, but are excreted rapidly by the kidneys. As a result, kidney toxicity is the main adverse effect, although ototoxicity and neuromuscular blockade can also occur. Gentamicin acts by interfering with bacterial protein synthesis. This compound is more potent than most other antibacterial inhibitors of protein synthesis, which are merely bacteriostatic, and its effects on the body are, likewise, more severe (Goodman & Gilman's The Pharmalogical Basis of Therapeutics 9th ed., pp. 1103-1115, J.G. Hardman et al. Eds., McGraw Hill, New York, 1996).

Aminoglycosides work rapidly, and the rate of bacterial killing is concentration-dependent. Residual bactericidal activity remains after serum concentration has fallen below the minimum inhibitory concentration (MIC), with a duration that is also dosage/concentration-dependent. The residual activity allows for once-a-day administration in some patients. These drugs diffuse into bacterial cells through porin channels in the outer membrane and are then transported across the cytoplasmic

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membrane via a membrane potential that is negative on the inside (Goodman & Gilman, supra).

Kidney damage, which can develop into renal failure, is due to the attack of gentamicin on the proximal convoluted tubule, particularly in the S1 and S2 segments. The necrosis, however, is often patchy and focal (Shanley *et al.* (1990), *Ren Fail* 12: 83-87). A rat study by Shanley *et al.* showed that superficial nephrons are more susceptible to necrosis than juxtamedullary nephrons, although the initial segment of the superficial nephrons is remarkably resistant to necrosis.

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Reported enzymatic changes upon gentamicin treatment are increased activities of N-acetyl-beta-D-glucosaminidase and alkaline phosphatase and decreased activities of sphingomyelinase, cathepsin B, Na⁺/K⁺-ATPase, lactate dehydrogenase and NADPH cytochrome C reductase, along with decreased protein synthesis and alpha-methylglucose transport (Monteil *et al.* (1993), *Ren Fail* 15: 475-483). An increase in gamma-glutamyl transpeptidase activity in urine has also been reported (Kocaoglu *et al.* (1994), *Arch Immunol Ther Exp (Warsz)* 42: 125-127), and the quantification of this enzyme in urine is a useful marker for monitoring gentamicin toxicity.

One source of renal pathology resulting from gentamicin treatment is the generation of reactive oxygen metabolites. Gentamicin has been shown, both *in vitro* and *in vivo*, to be capable of enhancing the production of reactive oxygen species. Iron, a necessary co-factor that catalyzes free-radical formation, is supplied by cytochrome P450 (Baliga *et al.* (1999), *Drug Metab Rev* 31: 971-997).

A gene delivery experiment in rats, in which the human kallikrein gene was cloned into an adenovirus vector and the construct then co-administered with a gentamicin preparation, showed that kallikrein can protect against gentamicin-induced nephrotoxicity. Significantly increased renal blood flow, glomerular filtration rates and urine flow were observed, along with decreased renal tubular damage, cellular necrosis and lumenal protein casts. Kallikrein gene delivery also caused a decrease in blood urea nitrogen levels and increases in urinary kinin and nitrite/nitrate levels. This study provides evidence that the tissue kallikrein-kinin system may be a key pathway that is perturbed during the induction of nephrotoxicity by gentamicin (Murakami *et al.* (1998), *Kidney Int* 53: 1305-1313).

Ifosfamide, an alkylating agent, is commonly used in chemotherapy to treat testicular, cervical, and lung cancer. Ifosfamide is slowly activated in the liver by hydroxylation, forming the triazene derivative 5-(3,3-dimethyl-1-triazeno)-imidazole-4-carboxamide (DTIC) (Goodman & Gilman's The Pharmacological Basis of Therapeutics 9th ed., p.1235, J.G. Hardman et al., Eds., McGraw Hill, New York, 1996). Cytochrome P450 activates DTIC via an N-demethylation reaction yielding an alkylating moiety, diazomethane. The active metabolites are then able to cross-link DNA causing growth arrest and cell death. Though ifosfamide is therapeutically useful, it is also associated with nephrotoxicity, urotoxicity, and central neurotoxicity.

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Mesna, another therapeutic, is often administered concomitantly to prevent kidney and bladder problems from arising (Brock and Pohl (1986), *IARC Sci Publ* 78: 269-279). However, there are documented cases in which tubular toxicity occurred and elevated urinary levels of alanine aminopeptidase and N-acetyl-beta-D-glucosaminidase were found in patients even though mesna was administered alongside ifosfamide (Goren *et al.* (1987), *Cancer Treat Rep* 71: 127-130).

One study examined 42 patients that had been administered ifosfamide to treat advanced soft-tissue sarcoma (Stuart-Harris *et al.* (1983), *Cancer Chemother Pharmacol* 11: 69-72). The ifosfamide dosage varied from 5.0 g/m² to 8.0 g/m², and all of the . patients were given mesna to counteract the negative effects of ifosfamide. Even so, nausea and vomiting were common to all of the patients. Out of the 42 patients, seven developed nephrotoxicity, and two of the cases progressed to fatal renal failure.

In another clinical study, renal tubular function was monitored in 18 neuroblastoma patients (Caron *et al.* (1992), *Med Pediatr Oncol* 20: 42-47). Tubular toxicity occurred in at least 12 of the patients, and seven of those patients eventually developed Debre-de Toni-Fanconi syndrome, although in 3 cases the syndrome was reversible.

Fanconi syndrome is a disorder marked by dysfunction of the proximal tubules of the kidney. It is associated with aminoaciduria, renal glycosuria, and hyperphosphaturia. Ifosfamide is often used experimentally on rats to induce Fanconi syndrome. In one study, rats that were administered 80 mg/kg of ifosfamide had significantly lower body weight and hematocrit than control rats (Springate and Van Liew (1995), *J Appl Toxicol* 15: 399-402). Additionally, the rats had low-grade glucosuria, proteinuria, and

phosphaturia. In a mouse study, ifosfamide induced elevated serum creatinine and urea levels and decreased the clearance rate of creatinine (Badary (1999), *J Ethnopharmacol* 67: 135-142).

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Cyclophosphamide, a nitrogen mustard and alkylating agent, is highly toxic to dividing cells and is commonly used in chemotherapy to treat malignant lymphomas, such as non-Hodgkin's lymphomas and Burkitt's lymphoma, multiple myeloma, leukemias, neuroblastomas, ovarian adenocarcinomas and retinoblastomas, as well as breast and lung cancer (Goodman & Gilman's The Pharmacological Basis of Therapeutics 9th ed., pp.1234, 1237-1239, J.G. Hardman et al., eds., McGraw Hill, New York, 1996; Physicians Desk Reference, 47th ed., pp. 744-745, Medical Economics Co., Inc., Montvale, NJ, 1993). Additionally, cyclophosphamide is used as an immunosuppressive agent in bone marrow transplantation and following organ transplantation. Although cyclophosphamide is therapeutically useful against certain types of cancer, it is also associated with cardiotoxicity, nephrotoxicity (including renal tubular necrosis), hemorrhagic cystitis, myelosuppression, hepatotoxicity, impairment of male and female reproductive systems, interstitial pneumonitis and central nervous system toxicity.

Once in the liver, cyclophosphamide is hydroxylated by the cytochrome P450 mixed function oxidase system, producing the active metabolites phosphoramide mustard and acrolein, which cross-link DNA and cause growth arrest and cell death. These metabolites, however, are highly toxic and cause adverse effects in the other organs into which they are transported, such as the kidneys. Acrolein is removed from the kidneys by secretion into the urine, resulting in cystitis (inflammation of the bladder), often hemorrhagic cystitis.

In the kidney, cyclophosphamide induces necrosis of the renal distal tubule. Cyclophosphamide, which is structurally similar to the anti-cancer drug ifosfamide, does not induce damage to the renal proximal tubule nor does it induce Debre-de Toni-Fanconi syndrome (Rossi *et al.* (1997), *Nephrol Dial Transplant* 12: 1091-1092).

One clinical trial of patients being treated with cyclophosphamide showed that renal damage from the drug leads to a reduced biotransformation rate and low renal clearance of the drug, resulting in a build-up of toxic alkylating metabolic products (Wagner *et al.* (1980), *Arzneimittelforschung* 30: 1588-1592).

In a study of patients suffering from malignant lymphomas and mammary carcinomas, a direct relationship was found between the dose of cyclophosphamide used in treatment and the concentration of alkylating metabolites in the patients' urine. The upper limit of the dose was determined by the nature and degree of the toxic side effects, rather than by the rate at which the drug could be metabolized (Saul *et al.* (1979), *J Cancer Res Clin Oncol* 94: 277-286). It is the acrolein itself that is toxic, not the alkylating activity of cyclophosphamide (Brock *et al.* (1979), *Arzneimittelforschung* 29: 659-661). A study on rats also showed that acrolein from the kidneys can produce hemorrhagic cystitis and that the acrolein concentration is directly related to the frequency and severity of the cystitis (Chijiwa *et al.* (1983), *Cancer Res* 43: 5205-5209).

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Carboplatin, a platinum coordination complex, is commonly used in chemotherapy as an anti-tumor agent. As a chemotherapeutic agent, carboplatin acts similarly to cisplatin. Carboplatin enters the cell by diffusion where it is activated by hydrolysis (Goodman & Gilman's The Pharmacological Basis of Therapeutics 9th ed., p. 1270-1271, J.G. Hardman et al. Eds., McGraw Hill, New York 1996). Once activated, the platinum complexes are able to react with DNA causing cross-linking to occur. One of the differences between carboplatin and cisplatin is that carboplatin is better tolerated clinically. Some of the side-effects associated with cisplatin, such as nausea, neurotoxicity, and nephrotoxicity, are seen at a lesser degree in patients administered carboplatin. Some other side-effects are hypomagnesaemia and hypokalaemia (Kintzel (2001), *Drug Saf* 24: 19-38).

In one study on male Wistar rats, carboplatin was administered at a dosage of 65mg/kg (Wolfgang *et al.* (1994), *Fundam Appl Toxicol* 22: 73-79). After treatment with carboplatin, CGT excretion was increased approximately two-fold.

Another study compared cisplatin and carboplatin when given in combination with vindesine and mitomycin C (Jelic *et al.* (2001) *Lung Cancer* 34: 1-13). The study showed that carboplatin administered with vindesine and mitomycin C was advantageous in terms of overall survival, although the regimen was more hematologically toxic than when cisplatin was given.

AY-25329, is a phenothiazine that has been shown to be mildly hepatotoxic and to induce nephrosis. Its structure is shown below.

Phenothiazines are a class of psychoactive drugs. They have been used to treat schizophrenia, paranoia, mania, hyperactivity in children, some forms of senility, and anxiety (http://www.encyclopedia.com/articlesnew/ 36591.html). Some side effects associated with prolonged use of the drugs are reduced blood pressure, Parkinsonism, reduction of motor activity, and visual impairment.

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Chlorpromazine (Thorazine or Largactil) is an aliphatic phenothiazine and is widely used for treating schizophrenia and manic depression. Prolactin secretion is increased while taking chlorpromazine, and galactorrhea and gynecomastia have both been associated with the drug (http://www.mentalhealth.com/drug/p30-c01.html). Trifluoperazine is another prescribed phenothiazine. It is used to treat anxiety, to prevent nausea and vomiting, and to manage psychotic disorders (http://www.mentalhealth.com/drug/p30-s04.html). Negative side-effects that have been associated with the drug are liver damage, bone marrow depression, and Parkinsonism.

Acyclovir (9-[(2-hydroxyethyl) methyl] guanine, Zovirax®), an anti-viral guanosine analogue, is used to treat herpes simplex virus (HSV), varicella zoster virus (VZV) and Epstein-Barr virus (EBV) infections. It is transported into cells by the nucleoside transporter that imports guanine, and acyclovir is phosphorylated by virally encoded thymidine kinase (TK). Other kinases convert acyclovir to its activated di- and triphosphate forms, which prevent the polymerization of viral DNA. Acyclovir triphosphate competes with dGTP for the viral polymerase, and acyclovir is preferentially incorporated, but as a monophosphate. As a result, chain elongation ceases (Fields Virology 3^d ed., Fields *et al.*, eds., pp. 436-440, Lippincott-Raven Publishers, Philadelphia, 1996; Cecil Textbook of Medicine, 20th ed., part XII, p. 1742, J. C. Bennett and F. Plum Eds., W. B. Saunders Co., Philadelphia, 1996).

The pharmacokinetics of acyclovir show that it has a useful half-life of about three hours and that most of it is excreted in the urine largely unchanged (Brigden *et al.* (1985), Scand J Infect Dis Suppl 47: 33-39). Not surprisingly, the most frequent adverse

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effect of acyclovir treatment is damage to various parts of the kidney, particularly the renal tubules. Crystalluria, or the precipitation of crystals (in this case, crystals of acyclovir), in the lumina of the renal tubules can occur (Fogazzi (1996), Nephrol Dial Transplant 11: 379-387). If the drug crystallizes in the renal collecting tubules, obstructive nephropathy and tubular necrosis can result (Richardson (2000), Vet Hum Toxicol 42: 370-371). Tissues from biopsies of affected patients showed dilation of the proximal and distal renal tubules, with loss of the brush border, flattening of the lining cells and focal nuclear loss (Becker et al. (1993), Am J Kidney Dis 22: 611-615).

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Citrinin, a mycotoxin produced by the fungus *Penicillium citrinum*, is a natural contaminant of foods and feeds (Bondy and Armstrong (1998) *Cell Biol. Toxicol.* 14: 323-332). It is known that mycotoxins can have negative effects on the immune system, however citrinin-treated animals have been shown to stimulate responses against antigens (Sharma (1993) *J. Dairy Sci.* 76: 892-897). Citrinin is a known nephrotoxin, and in birds such as chickens, ducklings, and turkeys, it causes diarrhea, increased food consumption and reduced weight gain due to kidney degeneration (Mehdi *et al.* (1981) *Food Cosmet. Toxicol.* 19: 723-733; Mehdi *et al.* (1984) *Vet. Pathol.* 21: 216-223). In the turkey and duckling study, both species exhibited nephrosis with the occurrence of hepatic and lymphoid lesions (Mehdi *et al.*, 1984).

In one study, citrinin was administered to rabbits as a single oral dose of either 120 or 67 mg/kg (Hanika et al. (1986) Vet. Pathol. 23: 245-253). Rabbits treated with citrinin exhibited renal alterations such as condensed and distorted mitochondria, distended intercellular spaces of the medullary and straight cortical distal tubules, and disorganization of interdigitating processes. In another rabbit study, citrininadministered rabbits displayed azotaemia and metabolic acidosis (Hanika et al. (1984) Food Chem. Toxicol. 22: 999-1008). Renal failure was indicated by decreased creatinine clearance and increased blood urea nitrogen and serum-creatinine levels.

In the past, mercury was an important component of pharmaceuticals, particularly of antiseptics, antibacterials, skin ointments, diuretics and laxatives. Although, mercury has been largely replaced by more effective, more specific and safer compounds, making drug-induced mercury poisoning rare, it is still widely used in industry. Poisoning from occupational exposure and environmental pollution, such as mercury release into public water supplies, remains a concern as wildlife, domestic animals and humans are affected.

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Because of their lipid solubility and ability to cross the blood-brain barrier, the most dangerous form of mercury is the organomercurials, the most common of which is methylmercury, a fungicide used for disinfecting crop seeds. In a number of countries, incidents involving large-scale illness and death from mercury poisoning have been reported when mercury-contaminated seeds were planted and the crops harvested and consumed. A second source of organic mercury poisoning results from industrial chemicals containing inorganic mercury, such as mercury catalysts, which form methylmercury as a reaction product. If this waste product is released into reservoirs, lakes, rivers or bays, the surrounding population can become sick or die, particularly those who eat local fish.

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The inorganic salt mercuric chloride, HgCl₂, as well as other mercuric salts, are more irritating and more toxic than the mercurous forms. Mercuric chloride is used today in industry, for the manufacture of bleach, electronics, plastics, fungicides and dental amalgams. The main source of human exposure is industrial dumping into rivers (Goodman & Gilman's: The Pharmacological Basis of Therapeutics (9th ed.), pp. 1654-1659, McGraw-Hill, New York, 1996).

When inorganic mercury salts are ingested, about 10% of the mercuric ions are absorbed by the gastrointenstinal tract, and a considerable portion of the Hg²⁺ can remain bound to the mucosal surfaces. The highest concentration of Hg²⁺ is found in the kidneys, as it is retained there longer than in other tissues. Consequently, the kidneys are the organ most adversely affected by inorganic mercury poisoning. The proximal tubules are the major site of damage, where tubular necrosis results. The mercury affects primarily the S2 and S3 portions of the proximal tubules, but, at high levels of mercury exposure, the S1 and distal portions of the tubules are also damaged. These regions of the nephrons are affected because they contain enzymes (such as gammaglutamyltranspeptidase) and transport proteins (such as the basolateral organic anion transport system) involved in mercury uptake (Diamond *et al.* (1998), *Toxicol Pathol* 26: 92-103).

Urinary markers of mercury toxicity which can be detected in NMR spectra include elevated levels of lactate, acetate and taurine and decreased levels of hippurate (Holmes *et al.* (2000), *Chem Res Toxicol* 13: 471-478). Known changes in gene expression in kidneys exposed to Hg²⁺ include up-regulation of the heat-shock protein

hsp72 and of the glucose-regulated protein grp94. The degree of tissue necrosis and level of expression of these proteins is proportional to both the dose of mercury (Hg²⁺) and the length of the exposure time to mercury (Hg²⁺), with hsp72 accumulating in the renal cortex and grp94 accumulating in the renal medulla (Goering *et al.* (2000), *Toxicol Sci* 53: 447-457).

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Diflunisal, a non-steroidal anti-inflammatory drug (NSAID), is a difluorophenyl derivative of salicylic acid (Goodman & Gilman's The Pharmacological Basis of Therapeutics 9th ed., p. 631, J.G. Hardman et al., Eds., McGraw Hill, New York, 1996). It is most frequently used in the treatment of osteoarthritis and musculoskeletal strains.

NSAIDs have analgesic, antipyretic and anti-inflammatory actions, however hepatotoxicity is known to be an adverse side effect of NSAID treatment (Masubuchi et al. (1998) J. Pharmacol. Exp. Ther. 287: 208-213). Diflunisal has been shown to be less toxic than other NSAIDs, nevertheless over long periods of dosage it can lead to deleterious effects on platelet or kidney function (Bergamo et al. (1989) Am. J. Nephrol.

9: 460-463). Other side effects that have been associated with diflunisal treatment are diarrhea, dizziness, drowsiness, gas or heartburn, headache, nausea, vomiting, and insomnia (http://arthritisinsight.com/medical/ meds/dolobid.html).

Masubuchi *et al.* compared the hepatotoxicity of 18 acidic NSAIDs. In the study, diflunisal (administered at a concentration of 500 μ M) was shown to increase LDH leakage in rat hepatocytes, a marker for cell injury, when compared to the control sample. In addition, treatment with diflunisal led to decreased intracellular ATP concentrations.

One study compared the effects of diflunisal and ibuprofen when given to patients over a two week period (Muncie and Nasrallah (1989) Clin. Ther. 11: 539-544). In both the ibuprofen and the diflunisal group, two patients complained of abdominal cramping. The study indicated that even during short-term usage some gastrointestinal effects may occur. The toxic dose used in this study was chosen as one that did not induce significant gastric ulceration in rats. The group of rats given the high dosage of diflunisal had increased concentrations of creatinine which is consistent with renal injury, although dehydration may also cause increases in creatinine concentration.

Cidofovir (Vistide®) is an antiviral cytosine analog used in the treatment of viral infections such as herpesvirus, adenovirus, papillomavirus, poxvirus and hepadnavirus

(Goodman & Gilman's The Pharmacological Basis of Therapeutics 9th ed., p. 1216, J.G. Hardman *et al.*, Eds., McGraw Hill, New York, 1996). It is also useful for the treatment of cytomegalovirus (CMV) infection, which is a type of herpesvirus.

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Some mild side effects seen in patients receiving cidofovir are nausea, vomiting, and fever. The most serious reported side effect of the drug is kidney toxicity (http://tthivclinic.com/cido.html). In response to the threat of nephrotoxicity, it is necessary for patients receiving cidofovir to have their kidneys checked before treatment, and the patients must be monitored during treatment for early symptoms of kidney problems. In addition, cidofovir is given with fluids to help reduce the risk of kidney toxicity (http://www.aidsinfonyc.org/ network/simple/cido.html). Probenecid, a drug that helps protect the kidneys, is normally administered concomitantly (Lalezari and Kuppermann (1997) *J. Acquir. Immune Defic. Syndr. Hum. Retrovirol.* 14: S27-31).

One study compared the safety and efficacy of cidofovir in the treatment of CMV (Lalezari et al. (1998) *J. Acquir. Immune Defic. Syndr. Hum. Retrovirol.* 17: 339-344). Approximately 40% of the patients exhibited dose-dependent asymptomatic proteinuria and 25% of the patients had elevated serum creatinine levels.

Pamidronate (Aredia®) is a bisphosphonate drug that is clinically used to inhibit bone resorption and make bones more stable. It is used to treat hypercalcemia (too much calcium in the blood) that occurs with some types of cancer. Typically administered by intravenous injection, pamidronate is frequently used in patients with breast cancer or multiple myeloma whose disease has spread to the bones. Some side effects related to pamidronate treatment are abdominal cramps, chills, confusion, fever, muscle spasms, nausea, muscle stiffness, and swelling at the injection site (http://www.nursing.uiowa.edu/sites/PedsPain/Adjuvants/PAMIDRnt.html). Patients with kidney problems may be prohibited from using pamidronate as it is excreted through the kidneys.

In one study, rats and mice were given varying doses of labeled pamidronate (Cal and Daley-Yates (1990) *Toxicology* 65: 179-197). Pamidronate treatment led to significant weight loss and a decrease in creatinine clearance. Morphological studies showed a loss of brush border membranes and the presence of focal proximal tubular necrosis.

Another study compared the tolerability of different treatments for hypercalcemia of malignancy by reviewing articles published between 1979 and 1998 (Zojer *et al.*

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(1999) *Drug Saf.* 21: 389-406). The authors found that elevated serum creatinine level, nausea, and fever were reported following treatment with bisphosphonates such as pamidronate.

Markowitz et al. (2001, J. Am. Soc. Nephrol. 12: 1164-1172) tried to determine whether there was a correlation between pamidronate treatment and collapsing focal segmental glomerulosclerosis (FSGS). The authors examined the histories of seven patients who had developed collapsing FSGS, and they found that the only drug treatment in common was the administration of pamidronate. When given at the recommended dose of 90 mg per month, renal toxicity was rare. However, when pamidronate was given at higher doses nephrotoxicity occurred.

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Lithium, an alkali metal, is the main pharmacological treatment for bipolar disorders. It is typically given as a salt, such as lithium carbonate or lithium citrate. Some common side effects of lithium treatment are an increase in urination, increase in drinking, dry mouth, weight gain, fine tremor, and fatigue. Some more serious side effects related to lithium treatment are blurred vision, mental confusion, seizures, vomiting, diarrhea, muscle weakness, drowsiness, and coarse tremor (Goodman & Gilman's The Pharmacological Basis of Therapeutics 9th ed., p. 448, J.G. Hardman et al., Eds., McGraw Hill, New York, 1996).

Since lithium is often used on a maintenance basis for a lifelong period, numerous studies have been performed to try and elucidate the effects of lithium on the kidney. One group administered lithium in daily doses within the human therapeutic range to male Wistar rats (Kling et al. (1984) Lab Invest 50: 526-535). Rats that were given lithium developed marked polyuria within three weeks of the initial dosing. The rats displayed elevated free water clearance and vasopressin-resistant diabetes insipidus. The cortical collecting tubules displayed morphological changes, e.g. dilation of the tubules, bulging cells lining the tubules, enlarged nuclei, following lithium treatment.

Another study examined a human population that had been given lithium for the treatment of bipolar disorder (Markowitz et al. (2000) J. Am. Soc. Nephrol. 11: 1439-1448). The patients had a mean age of 42.5 years and had been undergoing lithium treatment from 2 to 25 years (mean of 13.6 years). Approximately one fourth of the patients had nephrotic proteinuria, almost 90% of them had nephrogenic diabetes insipidus (NDI), and renal biopsies revealed a chronic tubulointerstitial nephropathy in

all of the patients. Following cessation of lithium treatment, seven of the patients proceeded to end-stage renal disease.

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Even though nephrotoxicity is a known side effect of lithium treatment, some studies have indicated that in actuality it is not all that common (Johnson (1998) Neuropsychopharmacology 19: 200-205). One study showed that the NDI-like effect in lithium treatment was easily overcome by increasing the levels of arginine vasopressin (AVP) (Carney et al. (1996) Kidney Int 50: 377-383). Other studies have suggested that patients with psychiatric disorders display certain defects in renal function without undergoing lithium treatment (Gitlin (1999) Drug Saf 20: 231-243).

Hydralazine, an antihypertensive drug, causes relaxation of arteriolar smooth muscle. Such vasodilation is linked to vigorous stimulation of the sympathetic nervous system, which in turn leads to increased heart rate and contractility, increased plasma renin activity, and fluid retention (Goodman & Gilman's The Pharmacological Basis of Therapeutics 9th ed., p. 794, J.G. Hardman *et al.*, Eds., McGraw Hill, New York, 1996). The increased renin activity leads to an increase in angiotensin II, which in turn causes stimulation of aldosterone and sodium reabsorption.

Hydralazine is used for the treatment of high blood pressure (hypertension) and for the treatment of pregnant women suffering from high blood pressure (pre-eclampsia or eclampsia). Some common side effects associated with hydralazine use are diarrhea, rapid heartbeat, headache, decreased appetite, and nausea. Hydralazine is often used concomitantly with drugs that inhibit sympathetic activity to combat the mild pulmonary hypertension that can be associated with hydralazine usage.

In one hydralazine study, rats were fed hydralazine and mineral metabolism was monitored (Peters *et al.* (1988) *Toxicol Lett* 41: 193-202). Manganese and zinc concentrations were not effected by hydralazine treatment, however tissue iron concentrations were decreased and kidney copper concentrations were increased compared to control groups.

Another study compared the effects of hydrazine, phenelzine, and hydralazine treatment on rats (Runge-Morris *et al.* (1996) *Drug Metab Dispos* 24: 734-737). Hydralazine caused an increase in renal GST-alpha subunit expression, although unlike hydrazine and phenelzine it did not alter renal cytochrome P4502E1 expression.

Colchicine, an alkoloid of *Colchicum autumale*, is an antiinflammatory agent used in the treatment of gouty arthritis (<u>Goodman & Gilman's The Pharmacological</u>

<u>Basis of Therapeutics 9th ed.</u>, p. 647, J.G. Hardman *et al.*, Eds., McGraw Hill, New York, 1996).

An antimitotic agent, colchicine binds to tubulin which leads to depolymerization and disappearance of the fibrillar microtubules in granulocytes and other motile cells. In doing so, the migration of granulocytes into the inflamed area is inhibited. Through a series of events, the inflammatory response is blocked.

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Some common, mild side effects associated with colchicine treatment are loss of appetite and hair loss. More severe side effects that warrant cessation of treatment are nausea, vomiting, diarrhea, and abdominal pain. Colchicine overdose can induce multiorgan failure with a high incidence of mortality. In this setting, renal failure is multifactorial and related to prolonged hypotension, hypoxemia, sepsis, and rhabdomyolysis. In rats, less dramatic doses have been shown to inhibit the secretion of many endogenous proteins such as insulin and parathyroid hormone.

One study investigated the effects of colchicine on microtubule polymerization status and post-translational modifications of tubulin in rat seminiferous tubules (Correa and Miller (2001) *Biol Reprod* 64: 1644-1652). Colchicine caused extensive microtubule depolymerization, and total tubulin levels decreased twofold after colchicine treatment. The authors also found that colchicine treatment led to a decrease in tyrosination of the microtubule pool of tubulin which was associated with depolymerization of microtubules.

Sulfadiazine, a sulfonamide, is an antimicrobial agent. It is commonly used concomitantly with pyrimethamine to treat toxoplasmosis, an infection of the brain, in patient suffering from AIDS. These drugs are able to cross the blood-brain barrier and are used at relatively high doses. In addition, sulfadiazine has been shown to be effective at preventing certain types of meningococcal diseases and in treating urinary tract infections.

Sulfonamides in general are structural analogs of para-aminobenzoic acid (PABA). Because they are competitive antagonists of PABA, sulfonamides are effective against bacteria that are required to utilize PABA for the synthesis of folic acid

(Goodman & Gilman's The Pharmacological Basis of Therapeutics 9th ed., p. 1058-1060, J.G. Hardman *et al.*, Eds., McGraw Hill, New York, 1996).

The main side effects associated with sulfadiazine treatment are fever and skin rashes. Decreases in white blood cells, red blood cells, and platelets, nausea, vomiting, and diarrhea are some other side effects that may result from sulfadiazine treatment. The most troublesome problem with this drug for HIV/AIDS patients is kidney toxicity. These patients tend to use these drugs for extended periods of time, which puts a constant strain on the kidneys. In addition, kidney stones tend to form in the bladder and ureter thereby blocking the flow of urine. Kidney damage may result, and if left untreated kidney failure may occur. Therefore, patients being treated with sulfadiazine are instructed to increase their fluid intake in order to prevent crystal formation in the kidneys.

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One case study examined four HIV-positive patients who had been given sulfadiazine to treat toxoplasmosis (Crespo *et al.* (2000) *Clin Nephrol* 54: 68-72). All four of the patients, one of whom was a previously healthy person, developed oliguria, abdominal pain, renal failure, and displayed multiple radiolucent renal calculi in echography. Following extensive hydration and alcalinization, the renal function of the patients returned to normal.

Adriamycin, known generically as doxorubicin, is an anthracycline antibiotic produced by the fungus *Streptomyces peucetius*. It is an anti-tumor drug used in the treatment of breast, ovarian, bladder, and lung cancers as well as non-Hodgkin's lymphoma, Hodgkin's disease and sarcoma (Goodman & Gilman's The Pharmacological Basis of Therapeutics 9th ed., p. 1264-1265, J.G. Hardman *et al.*, Eds., McGraw Hill, New York, 1996).

Adriamycin has tetracycline ring structures with the sugar daunosamine attached by glycosidic linkage. It is able to intercalate with DNA, it affects DNA and RNA synthesis, and it can interact with cell membranes and alter their functions. Typically the drug is cell-cycle specific for the S phase of cell division. By binding to the cancer cells' DNA and blocking topoisomerase II, cancer cells are unable to divide and grow.

Some common side effects associated with adriamycin treatment are fatigue, a drop in white blood cell, red blood cell, or platelet count, hair loss, skin discoloration, and watery eyes (www.cancerhelp.org.uk/help/default.asp?page=4025). More serious

side effects include myocardial toxicity, ulceration and necrosis of the colon, and development of a second cancer.

Because of its utility in fighting cancer, numerous studies have been performed in attempts to further understand the mechanisms and effects of adriamycin. In one study, investigators injected mice with a single dose of adriamycin (Chen *et al.* (1998) *Nephron* 78: 440-452). The mice exhibited signs of combined glomerular albuminuria and immunoglublinuria, progressively elevated levels of nitrite/nitrate in the urine, abnormal renal function, and other symptoms indicative of focal segmental glomerulosclerosis.

In another study, rats were given adriamycin and the effects on angiotensin converting enzyme (ACE) were monitored (Venkatesan *et al.* (1993) *Toxicology* 85: 137-148). The rats developed glomerular and tubular injury, and serum ACE levels were significantly elevated 20, 25, and 30 days post-treatment. A different study followed rabbits for up to one year that were treated with either adriamycin, nephrectomy, or combinations thereof (Gadeholt-Gothlin *et al.* (1995) *Urol Res* 23: 169-173). The rabbits that were treated with adriamycin exhibited signs of nephrotoxicity at relatively low doses.

Toxicity Prediction and Modeling

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The genes and gene expression information, gene expression profiles, as well as the portfolios and subsets of the genes provided in Tables 1-5, may be used to predict at 20 least one toxic effect, including the nephrotoxicity of a test or unknown compound. As used, herein, at least one toxic effect includes, but is not limited to, a detrimental change in the physiological status of a cell or organism. The response may be, but is not required to be, associated with a particular pathology, such as tissue necrosis. Accordingly, the toxic effect includes effects at the molecular and cellular level. 25 Nephrotoxicity is an effect as used herein and includes but is not limited to the pathologies of nephritis, kidney necrosis, glomerular and tubular injury, and focal segmental glomerulosclerosis. As used herein, a gene expression profile comprises any quantitative representation of the expression of at least one mRNA species in a cell sample or population and includes profiles made by various methods such as differential 30 display, PCR, hybridization analysis, etc.

In general, assays to predict the toxicity or nephrotoxicity of a test agent (or compound or multi-component composition) comprise the steps of exposing a cell

population to the test compound, assaying or measuring the level of relative or absolute gene expression of one or more of the genes in Tables 1-5 and comparing the identified expression level(s) to the expression levels disclosed in the Tables and database(s) disclosed herein. Assays may include the measurement of the expression levels of about 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25, 30, 50, 75, 100 or more genes from Tables 1-5.

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In the methods of the invention, the gene expression level for a gene or genes induced by the test agent, compound or compositions may be comparable to the levels found in the Tables or databases disclosed herein if the expression level varies within a factor of about 2, about 1.5 or about 1.0 fold. In some cases, the expression levels are comparable if the agent induces a change in the expression of a gene in the same direction (e.g., up or down) as a reference toxin.

The cell population that is exposed to the test agent, compound or composition may be exposed *in vitro* or *in vivo*. For instance, cultured or freshly isolated renal cells, in particular rat renal cells, may be exposed to the agent under standard laboratory and cell culture conditions. In another assay format, *in vivo* exposure may be accomplished by administration of the agent to a living animal, for instance a laboratory rat.

Procedures for designing and conducting toxicity tests in *in vitro* and *in vivo* systems are well known, and are described in many texts on the subject, such as Loomis *et al.*, Loomis's Esstentials of Toxicology, 4th Ed., Academic Press, New York, 1996; Echobichon, The Basics of Toxicity Testing, CRC Press, Boca Raton, 1992; Frazier, editor, *In Vitro* Toxicity Testing, Marcel Dekker, New York, 1992; and the like.

In *in vitro* toxicity testing, two groups of test organisms are usually employed:
One group serves as a control and the other group receives the test compound in a single dose (for acute toxicity tests) or a regimen of doses (for prolonged or chronic toxicity tests). Because, in some cases, the extraction of tissue as called for in the methods of the invention requires sacrificing the test animal, both the control group and the group receiving compound must be large enough to permit removal of animals for sampling tissues, if it is desired to observe the dynamics of gene expression through the duration of an experiment.

In setting up a toxicity study, extensive guidance is provided in the literature for selecting the appropriate test organism for the compound being tested, route of administration. dose ranges, and the like. Water or physiological saline (0.9% NaCl in

water) is the solute of choice for the test compound since these solvents permit administration by a variety of routes. When this is not possible because of solubility limitations, vegetable oils such as corn oil or organic solvents such as propylene glycol may be used.

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Regardless of the route of administration, the volume required to administer a given dose is limited by the size of the animal that is used. It is desirable to keep the volume of each dose uniform within and between groups of animals. When rats or mice are used, the volume administered by the oral route generally should not exceed about 0.005 ml per gram of animal. Even when aqueous or physiological saline solutions are used for parenteral injection the volumes that are tolerated are limited, although such solutions are ordinarily thought of as being innocuous. The intravenous LD₅₀ of distilled water in the mouse is approximately 0.044 ml per gram and that of isotonic saline is 0.068 ml per gram of mouse. In some instances, the route of administration to the test animal should be the same as, or as similar as possible to, the route of administration of the compound to man for therapeutic purposes.

When a compound is to be administered by inhalation, special techniques for generating test atmospheres are necessary. The methods usually involve aerosolization or nebulization of fluids containing the compound. If the agent to be tested is a fluid that has an appreciable vapor pressure, it may be administered by passing air through the solution under controlled temperature conditions. Under these conditions, dose is estimated from the volume of air inhaled per unit time, the temperature of the solution, and the vapor pressure of the agent involved. Gases are metered from reservoirs. When particles of a solution are to be administered, unless the particle size is less than about 2 µm the particles will not reach the terminal alveolar sacs in the lungs. A variety of apparatuses and chambers are available to perform studies for detecting effects of irritant or other toxic endpoints when they are administered by inhalation. The preferred method of administering an agent to animals is via the oral route, either by intubation or by incorporating the agent in the feed.

When the agent is exposed to cells *in vitro* or in cell culture, the cell population to be exposed to the agent may be divided into two or more subpopulations, for instance, by dividing the population into two or more identical aliquots. In some preferred embodiments of the methods of the invention, the cells to be exposed to the agent are

derived from kidney tissue. For instance, cultured or freshly isolated rat renal cells may be used.

The methods of the invention may be used generally to predict at least one toxic response, and, as described in the Examples, may be used to predict the likelihood that a compound or test agent will induce various specific kidney pathologies, such as nephritis, kidney necrosis, glomerular and tubular injury, focal segmental glomerulosclerosis, or other pathologies associated with at least one of the toxins herein described. The methods of the invention may also be used to determine the similarity of a toxic response to one or more individual compounds. In addition, the methods of the invention may be used to predict or elucidate the potential cellular pathways influenced, induced or modulated by the compound or test agent due to the similarity of the expression profile compared to the profile induced by a known toxin (see Tables 5-5CC).

Diagnostic Uses for the Toxicity Markers

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As described above, the genes and gene expression information or portfolios of the genes with their expression information as provided in Tables 1-5 may be used as diagnostic markers for the prediction or identification of the physiological state of tissue or cell sample that has been exposed to a compound or to identify or predict the toxic effects of a compound or agent. For instance, a tissue sample such as a sample of peripheral blood cells or some other easily obtainable tissue sample may be assayed by any of the methods described above, and the expression levels from a gene or genes from Tables 1-5 may be compared to the expression levels found in tissues or cells exposed to the toxins described herein. These methods may result in the diagnosis of a physiological state in the cell or may be used to identify the potential toxicity of a compound, for instance a new or unknown compound or agent. The comparison of expression data, as well as available sequence or other information may be done by researcher or diagnostician or may be done with the aid of a computer and databases as described below.

In another format, the levels of a gene(s) of Tables 1-5, its encoded protein(s), or any metabolite produced by the encoded protein may be monitored or detected in a sample, such as a bodily tissue or fluid sample to identify or diagnose a physiological state of an organism. Such samples may include any tissue or fluid sample, including urine, blood and easily obtainable cells such as peripheral lymphocytes.

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Use of the Markers for Monitoring Toxicity Progression

As described above, the genes and gene expression information provided in Tables 1-5 may also be used as markers for the monitoring of toxicity progression, such as that found after initial exposure to a drug, drug candidate, toxin, pollutant, etc. For instance, a tissue or cell sample may be assayed by any of the methods described above, and the expression levels from a gene or genes from Tables 1-5 may be compared to the expression levels found in tissue or cells exposed to the renal toxins described herein. The comparison of the expression data, as well as available sequence or other information may be done by a researcher or diagnostician or may be done with the aid of a computer and databases.

Use of the Toxicity Markers for Drug Screening

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According to the present invention, the genes identified in Tables 1-5 may be used as markers or drug targets to evaluate the effects of a candidate drug, chemical compound or other agent on a cell or tissue sample. The genes may also be used as drug targets to screen for agents that modulate their expression and/or activity. In various formats, a candidate drug or agent can be screened for the ability to stimulate the transcription or expression of a given marker or markers or to down-regulate or counteract the transcription or expression of a marker or markers. According to the present invention, one can also compare the specificity of a drug's effects by looking at the number of markers which the drug induces and comparing them. More specific drugs will have less transcriptional targets. Similar sets of markers identified for two drugs may indicate a similarity of effects.

Assays to monitor the expression of a marker or markers as defined in Tables 1-5 may utilize any available means of monitoring for changes in the expression level of the nucleic acids of the invention. As used herein, an agent is said to modulate the expression of a nucleic acid of the invention if it is capable of up- or down-regulating expression of the nucleic acid in a cell.

In one assay format, gene chips containing probes to one, two or more genes from Tables 1-5 may be used to directly monitor or detect changes in gene expression in the treated or exposed cell. Cell lines, tissues or other samples are first exposed to a test agent and in some instances, a known toxin, and the detected expression levels of one or more, or preferably 2 or more of the genes of Tables 1-5 are compared to the expression

levels of those same genes exposed to a known toxin alone. Compounds that modulate the expression patterns of the known toxin(s) would be expected to modulate potential toxic physiological effects *in vivo*. The genes in Tables 1-5 are particularly appropriate markers in these assays as they are differentially expressed in cells upon exposure to a known renal toxin. Tables 1 and 2 disclose those genes that are differentially expressed upon exposure to the named toxins and their corresponding GenBank Accession numbers. Table 3 discloses the human homologues and the corresponding GenBank Accession numbers of the differentially expressed genes of Tables 1 and 2.

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In another format, cell lines that contain reporter gene fusions between the open reading frame and/or the transcriptional regulatory regions of a gene in Tables 1-5 and any assayable fusion partner may be prepared. Numerous assayable fusion partners are known and readily available including the firefly luciferase gene and the gene encoding chloramphenical acetyltransferase (Alam *et al.* (1990), *Anal Biochem* 188: 245-254). Cell lines containing the reporter gene fusions are then exposed to the agent to be tested under appropriate conditions and time. Differential expression of the reporter gene between samples exposed to the agent and control samples identifies agents which modulate the expression of the nucleic acid.

Additional assay formats may be used to monitor the ability of the agent to modulate the expression of a gene identified in Tables 1-5. For instance, as described above, mRNA expression may be monitored directly by hybridization of probes to the nucleic acids of the invention. Cell lines are exposed to the agent to be tested under appropriate conditions and time, and total RNA or mRNA is isolated by standard procedures such those disclosed in Sambrook *et al.* (Molecular Cloning: A Laboratory Manual, 2nd Ed., Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY, 1989).

In another assay format, cells or cell lines are first identified which express the gene products of the invention physiologically. Cells and/or cell lines so identified would be expected to comprise the necessary cellular machinery such that the fidelity of modulation of the transcriptional apparatus is maintained with regard to exogenous contact of agent with appropriate surface transduction mechanisms and/or the cytosolic cascades. Further, such cells or cell lines may be transduced or transfected with an expression vehicle (e.g., a plasmid or viral vector) construct comprising an operable non-translated 5'-promoter containing end of the structural gene encoding the gene products

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of Tables 1-5 fused to one or more antigenic fragments or other detectable markers, which are peculiar to the instant gene products, wherein said fragments are under the transcriptional control of said promoter and are expressed as polypeptides whose molecular weight can be distinguished from the naturally occurring polypeptides or may further comprise an immunologically distinct or other detectable tag. Such a process is well known in the art (see Sambrook *et al.*, *supra*).

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Cells or cell lines transduced or transfected as outlined above are then contacted with agents under appropriate conditions; for example, the agent comprises a pharmaceutically acceptable excipient and is contacted with cells comprised in an aqueous physiological buffer such as phosphate buffered saline (PBS) at physiological pH, Eagles balanced salt solution (BSS) at physiological pH, PBS or BSS comprising serum or conditioned media comprising PBS or BSS and/or serum incubated at 37°C. Said conditions may be modulated as deemed necessary by one of skill in the art. Subsequent to contacting the cells with the agent, said cells are disrupted and the polypeptides of the lysate are fractionated such that a polypeptide fraction is pooled and contacted with an antibody to be further processed by immunological assay (e.g., ELISA, immunoprecipitation or Western blot). The pool of proteins isolated from the agentcontacted sample is then compared with the control samples (no exposure and exposure to a known toxin) where only the excipient is contacted with the cells and an increase or decrease in the immunologically generated signal from the agent-contacted sample compared to the control is used to distinguish the effectiveness and/or toxic effects of the agent.

Another embodiment of the present invention provides methods for identifying agents that modulate at least one activity of a protein(s) encoded by the genes in Tables 1-5. Such methods or assays may utilize any means of monitoring or detecting the desired activity.

In one format, the relative amounts of a protein (Tables 1-5) between a cell population that has been exposed to the agent to be tested compared to an un-exposed control cell population and a cell population exposed to a known toxin may be assayed. In this format, probes such as specific antibodies are used to monitor the differential expression of the protein in the different cell populations. Cell lines or populations are exposed to the agent to be tested under appropriate conditions and time. Cellular lysates

may be prepared from the exposed cell line or population and a control, unexposed cell line or population. The cellular lysates are then analyzed with the probe, such as a specific antibody.

Agents that are assayed in the above methods can be randomly selected or rationally selected or designed. As used herein, an agent is said to be randomly selected when the agent is chosen randomly without considering the specific sequences involved in the association of a protein of the invention alone or with its associated substrates, binding partners, etc. An example of randomly selected agents is the use a chemical library or a peptide combinatorial library, or a growth broth of an organism.

As used herein, an agent is said to be rationally selected or designed when the agent is chosen on a nonrandom basis which takes into account the sequence of the target site and/or its conformation in connection with the agent's action. Agents can be rationally selected or rationally designed by utilizing the peptide sequences that make up these sites. For example, a rationally selected peptide agent can be a peptide whose amino acid sequence is identical to or a derivative of any functional consensus site.

The agents of the present invention can be, as examples, peptides, small molecules, vitamin derivatives, as well as carbohydrates. Dominant negative proteins, DNAs encoding these proteins, antibodies to these proteins, peptide fragments of these proteins or mimics of these proteins may be introduced into cells to affect function. "Mimic" used herein refers to the modification of a region or several regions of a peptide molecule to provide a structure chemically different from the parent peptide but topographically and functionally similar to the parent peptide (see G.A. Grant in: Molecular Biology and Biotechnology, Meyers, ed., pp. 659-664, VCH Publishers, New York, 1995). A skilled artisan can readily recognize that there is no limit as to the structural nature of the agents of the present invention.

Nucleic Acid Assay Formats

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The genes identified as being differentially expressed upon exposure to a known renal toxin (Tables 1-5) may be used in a variety of nucleic acid detection assays to detect or quantify the expression level of a gene or multiple genes in a given sample. The genes described in Tables 1-5 may also be used in combination with one or more additional genes whose differential expression is associate with toxicity in a cell or tissue. In preferred embodiments, the genes in Tables 1-5 may be combined with one or

more of the genes described in prior and related applications 60/292,335; 60/297,523; 60/298,925; 60/303,810; 60/303,807; 60/303,808; 60/315,047; 60/324,928; 60/330,867; 60/330,462; 60/331,805; 60/336,144; 60/340,873; 60/357,843; 60/357,842; 60/357,844; 60/364,134; 60/370,206; 60/370,247; 60/370,144; 60/371,679; 60/372,794, 09/917,800 and 10/060,087 all of which are incorporated by reference on page 1 of this application.

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Any assay format to detect gene expression may be used. For example, traditional Northern blotting, dot or slot blot, nuclease protection, primer directed amplification, RT- PCR, semi- or quantitative PCR, branched-chain DNA and differential display methods may be used for detecting gene expression levels. Those methods are useful for some embodiments of the invention. In cases where smaller numbers of genes are detected, amplification based assays may be most efficient. Methods and assays of the invention, however, may be most efficiently designed with hybridization-based methods for detecting the expression of a large number of genes.

Any hybridization assay format may be used, including solution-based and solid support-based assay formats. Solid supports containing oligonucleotide probes for differentially expressed genes of the invention can be filters, polyvinyl chloride dishes, particles, beads, microparticles or silicon or glass based chips, etc. Such chips, wafers and hybridization methods are widely available, for example, those disclosed by Beattie (WO 95/11755).

Any solid surface to which oligonucleotides can be bound, either directly or indirectly, either covalently or non-covalently, can be used. A preferred solid support is a high density array or DNA chip. These contain a particular oligonucleotide probe in a predetermined location on the array. Each predetermined location may contain more than one molecule of the probe, but each molecule within the predetermined location has an identical sequence. Such predetermined locations are termed features. There may be, for example, from 2, 10, 100, 1000 to 10,000, 100,000 or 400,000 or more of such features on a single solid support. The solid support, or the area within which the probes are attached may be on the order of about a square centimeter. Probes corresponding to the genes of Tables 1-5 or from the related applications described above may be attached to single or multiple solid support structures, *e.g.*, the probes may be attached to a single chip or to multiple chips to comprise a chip set.

Oligonucleotide probe arrays for expression monitoring can be made and used

according to any techniques known in the art (see for example, Lockhart *et al.* (1996), *Nat Biotechnol* 14: 1675-1680; McGall *et al.* (1996), *Proc Nat Acad Sci* USA 93: 13555-13460). Such probe arrays may contain at least two or more oligonucleotides that are complementary to or hybridize to two or more of the genes described in Tables 1-5. For instance, such arrays may contain oligonucleotides that are complementary to or hybridize to at least 2, 3, 4, 5, 6, 7, 8, 9, 10, 20, 30, 50, 70, 100 or more of the genes described herein. Preferred arrays contain all or nearly all of the genes listed in Tables 1-5, or individually, the gene sets of Tables 5-5CC. In a preferred embodiment, arrays are constructed that contain oligonucleotides to detect all or nearly all of the genes in any one of or all of Tables 1-5 on a single solid support substrate, such as a chip.

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The sequences of the expression marker genes of Tables 1-5 are in the public databases. Table 1 provides the GenBank Accession Number or NCBI RefSeq ID for each of the sequences (see www.ncbi.nlm.nih.gow). Table 3 provides the LocusLink and Unigene names and descriptions for the human homologues of the genes described in Tables 1 and 2. The sequences of the genes in GenBank and/or RefSeq are expressly herein incorporated by reference in their entirety as of the filing date of this application, as are related sequences, for instance, sequences from the same gene of different lengths, variant sequences, polymorphic sequences, genomic sequences of the genes and related sequences from different species, including the human counterparts, where appropriate. These sequences may be used in the methods of the invention or may be used to produce the probes and arrays of the invention. In some embodiments, the genes in Tables 1-5 that correspond to the genes or fragments previously associated with a toxic response may be excluded from the Tables.

As described above, in addition to the sequences of the GenBank Accession Numbers or NCBI RefSeq ID's disclosed in the Tables 1-5, sequences such as naturally occurring variants or polymorphic sequences may be used in the methods and compositions of the invention. For instance, expression levels of various allelic or homologous forms of a gene disclosed in Tables 1-5 may be assayed. Any and all nucleotide variations that do not alter the functional activity of a gene listed in the Tables 1-5, including all naturally occurring allelic variants of the genes herein disclosed, may be used in the methods and to make the compositions (e.g., arrays) of the invention.

Probes based on the sequences of the genes described above may be prepared by

any commonly available method. Oligonucleotide probes for screening or assaying a tissue or cell sample are preferably of sufficient length to specifically hybridize only to appropriate, complementary genes or transcripts. Typically the oligonucleotide probes will be at least about 10, 12, 14, 16, 18, 20 or 25 nucleotides in length. In some cases, longer probes of at least 30, 40, or 50 nucleotides will be desirable.

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As used herein, oligonucleotide sequences that are complementary to one or more of the genes described in Tables 1-5 refer to oligonucleotides that are capable of hybridizing under stringent conditions to at least part of the nucleotide sequences of said genes. Such hybridizable oligonucleotides will typically exhibit at least about 75% sequence identity at the nucleotide level to said genes, preferably about 80% or 85% sequence identity or more preferably about 90% or 95% or more sequence identity to said genes.

"Bind(s) substantially" refers to complementary hybridization between a probe nucleic acid and a target nucleic acid and embraces minor mismatches that can be accommodated by reducing the stringency of the hybridization media to achieve the desired detection of the target polynucleotide sequence.

The terms "background" or "background signal intensity" refer to hybridization signals resulting from non-specific binding, or other interactions, between the labeled target nucleic acids and components of the oligonucleotide array (e.g., the oligonucleotide probes, control probes, the array substrate, etc.). Background signals may also be produced by intrinsic fluorescence of the array components themselves. A single background signal can be calculated for the entire array, or a different background signal may be calculated for each target nucleic acid. In a preferred embodiment, background is calculated as the average hybridization signal intensity for the lowest 5% to 10% of the probes in the array, or, where a different background signal is calculated for each target gene, for the lowest 5% to 10% of the probes for each gene. Of course, one of skill in the art will appreciate that where the probes to a particular gene hybridize well and thus appear to be specifically binding to a target sequence, they should not be used in a background signal calculation. Alternatively, background may be calculated as the average hybridization signal intensity produced by hybridization to probes that are not complementary to any sequence found in the sample (e.g. probes directed to nucleic acids of the opposite sense or to genes not found in the sample such as bacterial genes

where the sample is mammalian nucleic acids). Background can also be calculated as the average signal intensity produced by regions of the array that lack any probes at all.

The phrase "hybridizing specifically to" or "specifically hybridizes" refers to the binding, duplexing, or hybridizing of a molecule substantially to or only to a particular nucleotide sequence or sequences under stringent conditions when that sequence is present in a complex mixture (e.g., total cellular) DNA or RNA.

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Assays and methods of the invention may utilize available formats to simultaneously screen at least about 100, preferably about 1000, more preferably about 10,000 and most preferably about 1,000,000 different nucleic acid hybridizations.

As used herein a "probe" is defined as a nucleic acid, capable of binding to a target nucleic acid of complementary sequence through one or more types of chemical bonds, usually through complementary base pairing, usually through hydrogen bond formation. As used herein, a probe may include natural (*i.e.*, A, G, U, C, or T) or modified bases (7-deazaguanosine, inosine, *etc.*). In addition, the bases in probes may be joined by a linkage other than a phosphodiester bond, so long as it does not interfere with hybridization. Thus, probes may be peptide nucleic acids in which the constituent bases are joined by peptide bonds rather than phosphodiester linkages.

The term "perfect match probe" refers to a probe that has a sequence that is perfectly complementary to a particular target sequence. The test probe is typically perfectly complementary to a portion (subsequence) of the target sequence. The perfect match (PM) probe can be a "test probe", a "normalization control" probe, an expression level control probe and the like. A perfect match control or perfect match probe is, however, distinguished from a "mismatch control" or "mismatch probe."

The terms "mismatch control" or "mismatch probe" refer to a probe whose sequence is deliberately selected not to be perfectly complementary to a particular target sequence. For each mismatch (MM) control in a high-density array there typically exists a corresponding perfect match (PM) probe that is perfectly complementary to the same particular target sequence. The mismatch may comprise one or more bases.

While the mismatch(es) may be located anywhere in the mismatch probe, terminal mismatches are less desirable as a terminal mismatch is less likely to prevent hybridization of the target sequence. In a particularly preferred embodiment, the mismatch is located at or near the center of the probe such that the mismatch is most

likely to destabilize the duplex with the target sequence under the test hybridization conditions.

The term "stringent conditions" refers to conditions under which a probe will hybridize to its target subsequence, but with only insubstantial hybridization to other sequences or to other sequences such that the difference may be identified. Stringent conditions are sequence-dependent and will be different in different circumstances. Longer sequences hybridize specifically at higher temperatures. Generally, stringent conditions are selected to be about 5°C lower than the thermal melting point (Tm) for the specific sequence at a defined ionic strength and pH.

Typically, stringent conditions will be those in which the salt concentration is at least about 0.01 to 1.0 M Na⁺ ion concentration (or other salts) at pH 7.0 to 8.3 and the temperature is at least about 30°C for short probes (*e.g.*, 10 to 50 nucleotides). Stringent conditions may also be achieved with the addition of destabilizing agents such as formamide.

The "percentage of sequence identity" or "sequence identity" is determined by comparing two optimally aligned sequences or subsequences over a comparison window or span, wherein the portion of the polynucleotide sequence in the comparison window may optionally comprise additions or deletions (*i.e.*, gaps) as compared to the reference sequence (which does not comprise additions or deletions) for optimal alignment of the two sequences. The percentage is calculated by determining the number of positions at which the identical submit (*e.g.* nucleic acid base or amino acid residue) occurs in both sequences to yield the number of matched positions, dividing the number of matched positions by the total number of positions in the window of comparison and multiplying the result by 100 to yield the percentage of sequence identity. Percentage sequence identity when calculated using the programs GAP or BESTFIT (see below) is calculated using default gap weights.

Probe design

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One of skill in the art will appreciate that an enormous number of array designs are suitable for the practice of this invention. The high density array will typically include a number of test probes that specifically hybridize to the sequences of interest. Probes may be produced from any region of the genes identified in the Tables and the attached representative sequence listing. In instances where the gene reference in the

Tables is an EST, probes may be designed from that sequence or from other regions of the corresponding full-length transcript that may be available in any of the sequence databases, such as those herein described. See WO 99/32660 for methods of producing probes for a given gene or genes. In addition, any available software may be used to produce specific probe sequences, including, for instance, software available from Molecular Biology Insights, Olympus Optical Co. and Biosoft International. In a preferred embodiment, the array will also include one or more control probes.

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High density array chips of the invention include "test probes." Test probes may be oligonucleotides that range from about 5 to about 500, or about 7 to about 50 nucleotides, more preferably from about 10 to about 40 nucleotides and most preferably from about 15 to about 35 nucleotides in length. In other particularly preferred embodiments, the probes are 20 or 25 nucleotides in length. In another preferred embodiment, test probes are double or single strand DNA sequences such as cDNA fragments. DNA sequences are isolated or cloned from natural sources or amplified from natural sources using native nucleic acid as templates. These probes have sequences complementary to particular subsequences of the genes whose expression they are designed to detect. Thus, the test probes are capable of specifically hybridizing to the target nucleic acid they are to detect.

In addition to test probes that bind the target nucleic acid(s) of interest, the high density array can contain a number of control probes. The control probes may fall into three categories referred to herein as 1) normalization controls; 2) expression level controls; and 3) mismatch controls.

Normalization controls are oligonucleotide or other nucleic acid probes that are complementary to labeled reference oligonucleotides or other nucleic acid sequences that are added to the nucleic acid sample to be screened. The signals obtained from the normalization controls after hybridization provide a control for variations in hybridization conditions, label intensity, "reading" efficiency and other factors that may cause the signal of a perfect hybridization to vary between arrays. In a preferred embodiment, signals (e.g., fluorescence intensity) read from all other probes in the array are divided by the signal (e.g., fluorescence intensity) from the control probes thereby normalizing the measurements.

Virtually any probe may serve as a normalization control. However, it is

recognized that hybridization efficiency varies with base composition and probe length. Preferred normalization probes are selected to reflect the average length of the other probes present in the array, however, they can be selected to cover a range of lengths. The normalization control(s) can also be selected to reflect the (average) base composition of the other probes in the array, however in a preferred embodiment, only one or a few probes are used and they are selected such that they hybridize well (i.e., no secondary structure) and do not match any target-specific probes.

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Expression level controls are probes that hybridize specifically with constitutively expressed genes in the biological sample. Virtually any constitutively expressed gene provides a suitable target for expression level controls. Typically expression level control probes have sequences complementary to subsequences of constitutively expressed "housekeeping genes" including, but not limited to the actin gene, the transferrin receptor gene, the GAPDH gene, and the like.

Mismatch controls may also be provided for the probes to the target genes, for expression level controls or for normalization controls. Mismatch controls are oligonucleotide probes or other nucleic acid probes identical to their corresponding test or control probes except for the presence of one or more mismatched bases. A mismatched base is a base selected so that it is not complementary to the corresponding base in the target sequence to which the probe would otherwise specifically hybridize. 20 One or more mismatches are selected such that under appropriate hybridization conditions (e.g., stringent conditions) the test or control probe would be expected to hybridize with its target sequence, but the mismatch probe would not hybridize (or would hybridize to a significantly lesser extent). Preferred mismatch probes contain a central mismatch. Thus, for example, where a probe is a 20 mer, a corresponding mismatch 25 probe will have the identical sequence except for a single base mismatch (e.g., substituting a G, a C or a T for an A) at any of positions 6 through 14 (the central mismatch).

Mismatch probes thus provide a control for non-specific binding or cross hybridization to a nucleic acid in the sample other than the target to which the probe is directed. For example, if the target is present the perfect match probes should be consistently brighter than the mismatch probes. In addition, if all central mismatches are present, the mismatch probes can be used to detect a mutation, for instance, a mutation of

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a gene in the accompanying Tables 1-5. The difference in intensity between the perfect match and the mismatch probe provides a good measure of the concentration of the hybridized material.

Nucleic Acid Samples

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Cell or tissue samples may be exposed to the test agent *in vitro* or *in vivo*. When cultured cells or tissues are used, appropriate mammalian cell extracts, such as liver extracts, may also be added with the test agent to evaluate agents that may require biotransformation to exhibit toxicity. In a preferred format, primary isolates of animal or human renal cells which already express the appropriate complement of drugmetabolizing enzymes may be exposed to the test agent without the addition of mammalian kidney extracts.

The genes which are assayed according to the present invention are typically in the form of mRNA or reverse transcribed mRNA. The genes may or may not be cloned. The genes may or may not be amplified. The cloning and/or amplification do not appear to bias the representation of genes within a population. In some assays, it may be preferable, however, to use polyA+ RNA as a source, as it can be used with less processing steps.

As is apparent to one of ordinary skill in the art, nucleic acid samples used in the methods and assays of the invention may be prepared by any available method or process. Methods of isolating total mRNA are well known to those of skill in the art. For example, methods of isolation and purification of nucleic acids are described in detail in Chapter 3 of Laboratory Techniques in Biochemistry and Molecular Biology, Vol. 24, Hybridization With Nucleic Acid Probes: Theory and Nucleic Acid Probes, P. Tijssen, Ed., Elsevier Press, New York, 1993. Such samples include RNA samples, but also include cDNA synthesized from a mRNA sample isolated from a cell or tissue of interest. Such samples also include DNA amplified from the cDNA, and RNA transcribed from the amplified DNA. One of skill in the art would appreciate that it is desirable to inhibit or destroy RNase present in homogenates before homogenates are used.

Biological samples may be of any biological tissue or fluid or cells from any organism as well as cells raised *in vitro*, such as cell lines and tissue culture cells. Frequently the sample will be a tissue or cell sample that has been exposed to a

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compound, agent, drug, pharmaceutical composition, potential environmental pollutant or other composition. In some formats, the sample will be a "clinical sample" which is a sample derived from a patient. Typical clinical samples include, but are not limited to, sputum, blood, blood-cells (e.g., white cells), tissue or fine needle biopsy samples, urine, peritoneal fluid, and pleural fluid, or cells therefrom. Biological samples may also include sections of tissues, such as frozen sections or formalin fixed sections taken for histological purposes.

Forming High Density Arrays

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Methods of forming high density arrays of oligonucleotides with a minimal number of synthetic steps are known. The oligonucleotide analogue array can be synthesized on a single or on multiple solid substrates by a variety of methods, including, but not limited to, light-directed chemical coupling, and mechanically directed coupling (see Pirrung, U.S. Patent No. 5,143,854).

In brief, the light-directed combinatorial synthesis of oligonucleotide arrays on a glass surface proceeds using automated phosphoramidite chemistry and chip masking techniques. In one specific implementation, a glass surface is derivatized with a silane reagent containing a functional group, *e.g.*, a hydroxyl or amine group blocked by a photolabile protecting group. Photolysis through a photolithographic mask is used selectively to expose functional groups which are then ready to react with incoming 5' photoprotected nucleoside phosphoramidites. The phosphoramidites react only with those sites which are illuminated (and thus exposed by removal of the photolabile blocking group). Thus, the phosphoramidites only add to those areas selectively exposed from the preceding step. These steps are repeated until the desired array of sequences have been synthesized on the solid surface. Combinatorial synthesis of different oligonucleotide analogues at different locations on the array is determined by the pattern of illumination during synthesis and the order of addition of coupling reagents.

In addition to the foregoing, additional methods which can be used to generate an array of oligonucleotides on a single substrate are described in PCT Publication Nos. WO 93/09668 and WO 01/23614. High density nucleic acid arrays can also be fabricated by depositing pre-made or natural nucleic acids in predetermined positions. Synthesized or natural nucleic acids are deposited on specific locations of a substrate by light directed targeting and oligonucleotide directed targeting. Another embodiment uses

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a dispenser that moves from region to region to deposit nucleic acids in specific spots.

Hybridization

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Nucleic acid hybridization simply involves contacting a probe and target nucleic acid under conditions where the probe and its complementary target can form stable hybrid duplexes through complementary base pairing. See WO 99/32660. The nucleic acids that do not form hybrid duplexes are then washed away leaving the hybridized nucleic acids to be detected, typically through detection of an attached detectable label. It is generally recognized that nucleic acids are denatured by increasing the temperature or decreasing the salt concentration of the buffer containing the nucleic acids. Under low stringency conditions (e.g., low temperature and/or high salt) hybrid duplexes (e.g., DNA:DNA, RNA:RNA, or RNA:DNA) will form even where the annealed sequences are not perfectly complementary. Thus, specificity of hybridization is reduced at lower stringency. Conversely, at higher stringency (e.g., higher temperature or lower salt) successful hybridization tolerates fewer mismatches. One of skill in the art will appreciate that hybridization conditions may be selected to provide any degree of stringency.

In a preferred embodiment, hybridization is performed at low stringency, in this case in 6x SSPET at 37°C (0.005% Triton X-100), to ensure hybridization and then subsequent washes are performed at higher stringency (e.g., 1x SSPET at 37°C) to eliminate mismatched hybrid duplexes. Successive washes may be performed at increasingly higher stringency (e.g., down to as low as 0.25x SSPET at 37°C to 50°C) until a desired level of hybridization specificity is obtained. Stringency can also be increased by addition of agents such as formamide. Hybridization specificity may be evaluated by comparison of hybridization to the test probes with hybridization to the various controls that can be present (e.g., expression level control, normalization control, mismatch controls, etc.).

In general, there is a tradeoff between hybridization specificity (stringency) and signal intensity. Thus, in a preferred embodiment, the wash is performed at the highest stringency that produces consistent results and that provides a signal intensity greater than approximately 10% of the background intensity. Thus, in a preferred embodiment, the hybridized array may be washed at successively higher stringency solutions and read between each wash. Analysis of the data sets thus produced will reveal a wash

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stringency above which the hybridization pattern is not appreciably altered and which provides adequate signal for the particular oligonucleotide probes of interest.

Signal Detection

The hybridized nucleic acids are typically detected by detecting one or more labels attached to the sample nucleic acids. The labels may be incorporated by any of a number of means well known to those of skill in the art. See WO 99/32660.

Databases

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The present invention includes relational databases containing sequence information, for instance, for the genes of Tables 1-5, as well as gene expression information from tissue or cells exposed to various standard toxins, such as those herein described (see Tables 5-5CC). Databases may also contain information associated with a given sequence or tissue sample such as descriptive information about the gene associated with the sequence information (see Tables 1 and 2), or descriptive information concerning the clinical status of the tissue sample, or the animal from which the sample was derived. The database may be designed to include different parts, for instance a sequence database and a gene expression database. Methods for the configuration and construction of such databases and computer-readable media to which such databases are saved are widely available, for instance, see U.S. Patent No. 5,953,727, which is herein incorporated by reference in its entirety.

The databases of the invention may be linked to an outside or external database such as GenBank (www.ncbi.nlm.nih.gov/entrez.index.html); KEGG (www.genome.ad.jp/kegg); SPAD (www.grt.kyushu-u.ac.jp/spad/index.html); HUGO (www.gene.ucl.ac.uk/hugo); Swiss-Prot (www.expasy.ch.sprot); Prosite (www.expasy.ch/tools/scnpsit1.html); OMIM (www.ncbi.nlm.nih.gov/omim); and GDB (www.gdb.org). In a preferred embodiment, as described in Tables 1-5, the external database is GenBank and the associated databases maintained by the National Center for Biotechnology Information (NCBI) (www.ncbi.nlm.nih.gov).

Any appropriate computer platform, user interface, etc. may be used to perform the necessary comparisons between sequence information, gene expression information and any other information in the database or information provided as an input. For example, a large number of computer workstations are available from a variety of manufacturers, such has those available from Silicon Graphics. Client/server

environments, database servers and networks are also widely available and appropriate platforms for the databases of the invention.

The databases of the invention may be used to produce, among other things, electronic Northerns that allow the user to determine the cell type or tissue in which a given gene is expressed and to allow determination of the abundance or expression level of a given gene in a particular tissue or cell.

The databases of the invention may also be used to present information identifying the expression level in a tissue or cell of a set of genes comprising one or more of the genes in Tables 1-5, comprising the step of comparing the expression level of at least one gene in Tables 1-5 in a cell or tissue exposed to a test agent to the level of expression of the gene in the database. Such methods may be used to predict the toxic potential of a given compound by comparing the level of expression of a gene or genes in Tables 1-5 from a tissue or cell sample exposed to the test agent to the expression levels found in a control tissue or cell samples exposed to a standard toxin or renal toxin such as those herein described. Such methods may also be used in the drug or agent screening assays as described herein.

Kits

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The invention further includes kits combining, in different combinations, high-density oligonucleotide arrays, reagents for use with the arrays, protein reagents encoded by the genes of the Tables, signal detection and array-processing instruments, gene expression databases and analysis and database management software described above. The kits may be used, for example, to predict or model the toxic response of a test compound, to monitor the progression of renal disease states, to identify genes that show promise as new drug targets and to screen known and newly designed drugs as discussed above.

The databases packaged with the kits are a compilation of expression patterns from human or laboratory animal genes and gene fragments (corresponding to the genes of Tables 1-5). In particular, the database software and packaged information that may contain the databases saved to a computer-readable medium include the expression results of Tables 1-5 that can be used to predict toxicity of a test agent by comparing the expression levels of the genes of Tables 1-5 induced by the test agent to the expression levels presented in Tables 5-5CC. In another format, database and software information

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may be provided in a remote electronic format, such as a website, the address of which may be packaged in the kit.

The kits may used in the pharmaceutical industry, where the need for early drug testing is strong due to the high costs associated with drug development, but where bioinformatics, in particular gene expression informatics, is still lacking. These kits will reduce the costs, time and risks associated with traditional new drug screening using cell cultures and laboratory animals. The results of large-scale drug screening of pre-grouped patient populations, pharmacogenomics testing, can also be applied to select drugs with greater efficacy and fewer side-effects. The kits may also be used by smaller biotechnology companies and research institutes who do not have the facilities for performing such large-scale testing themselves.

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Databases and software designed for use with microarrays is discussed in Balaban et al., U.S. Patent Nos. 6,229,911, a computer-implemented method for managing information, stored as indexed Tables 1-5, collected from small or large numbers of microarrays, and 6,185,561, a computer-based method with data mining capability for collecting gene expression level data, adding additional attributes and reformatting the data to produce answers to various queries. Chee et al., U.S. Patent No. 5,974,164, disclose a software-based method for identifying mutations in a nucleic acid sequence based on differences in probe fluorescence intensities between wild type and mutant sequences that hybridize to reference sequences.

Without further description, it is believed that one of ordinary skill in the art can, using the preceding description and the following illustrative examples, make and utilize the compounds of the present invention and practice the claimed methods. The following working examples therefore, specifically point out the preferred embodiments of the present invention, and are not to be construed as limiting in any way the remainder of the disclosure.

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EXAMPLES

Example 1: Identification of Toxicity Markers

The renal toxins cephaloridine, cisplatin, puromycin aminonucleoside (PAN), bromoethylamine hydrobromide (BEA), gentamicin, ifosfamide, cyclophosphamide, carboplatin, AY-25329, indomethacin, acyclovir, citrinin, mercuric chloride, diflunisal, cidofovir, pamidronate, lithium, hydralazine, colchicine, sulfadiazine, and adriamycin and control compositions were administered to male Sprague-Dawley rats at various timepoints using administration diluents, protocols and dosing regimes as previously described in the art and previously described in the priority applications discussed above.

10 The low and high dose level for each compound are provided in the chart below.

Renal Toxin	Low Dose (mg/kg)	High Dose (mg/kg)	Method of Administration
cephaloridine	100	800	intravenous
cisplatin	1	5	intravenous
PAN	10	150	intravenous
BEA	10	200	intraperitoneal
gentamicin	2	80	intramuscular
ifosfamide	5	100	intraperitoneal
cyclophosphamide	20	2000	intraperitoneal
carboplatin	5	50	intravenous
AY-25329	25	250	oral gavage
indomethacin	1	10	oral gavage
acyclovir	10	100	intraperitoneal
citrinin	1	35	intraperitoneal
mercuric chloride	0.1	1	intravenous
diflunisal	2	400	oral gavage
cidofovir	10	100	intraperitoneal
pamidronate	1	60	intraperitoneal
lithium	0.3 (nmol/kg)	3 (nmol/kg)	intraperitoneal
hydralazine	2.5	25	intraperitoneal
colchicine	0.15	1.5	intraperitoneal
sulfadiazine	100	1000	intravenous
adriamycin	1.3	12.8	intravenous

After administration, the dosed animals were observed and tissues were collected as described below:

OBSERVATION OF ANIMALS

1. Clinical Observations-

Twice daily: mortality and moribundity check.

Cage Side Observations - skin and fur, eyes and
mucous membrane, respiratory system, circulatory
system, autonomic and central nervous system,
somatomotor pattern, and behavior pattern.

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Potential signs of toxicity, including tremors, convulsions, salivation, diarrhea, lethargy, coma or other atypical behavior or appearance, were recorded as they occurred and included a time of onset, degree, and duration.

Prior to randomization, prior to initial treatment,

2. Physical Examinations-

Prior to randomization, prior to initial treatment, and prior to sacrifice.

3. Body Weights-

Prior to randomization, prior to initial treatment, and prior to sacrifice.

10 CLINICAL PATHOLOGY

1. Frequency Prior to necropsy.

2. Number of animals

All surviving animals.

3. Bleeding Procedure

Blood was obtained by puncture of the orbital sinus while under 70% $CO_2/30\%$ O_2 anesthesia.

 Collection of Blood Samples Approximately 0.5 mL of blood was collected into EDTA tubes for evaluation of hematology parameters. Approximately 1 mL of blood was collected into serum separator tubes for clinical chemistry analysis. Approximately 200 uL of plasma was obtained and frozen at ~-80°C for

test compound/metabolite estimation. An

additional ~2 mL of blood was collected into a 15 mL conical polypropylene vial to which ~3 mL of Trizol was immediately added. The contents were immediately mixed with a vortex and by repeated inversion. The tubes were frozen in liquid nitrogen

and stored at ~--80°C.

TERMINATION PROCEDURES

Terminal Sacrifice

Approximately 3, 6, 24, 48, 72, 120, 144, 168, 336, and/or 360 hours after

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the initial dose, rats were weighed, physically examined, sacrificed by decapitation, and exsanguinated. The animals were necropsied within approximately five minutes of sacrifice. Separate sterile, disposable instruments were used for each animal, with the exception of bone cutters, which were used to open the skull cap. The bone cutters were dipped in disinfectant solution between animals.

Necropsies were conducted on each animal following procedures approved by board-certified pathologists.

Animals not surviving until terminal sacrifice were discarded without necropsy (following euthanasia by carbon dioxide asphyxiation, if moribund). The approximate time of death for moribund or found dead animals was recorded.

Postmortem Procedures

Fresh and sterile disposable instruments were used to collect tissues. Gloves were worn at all times when handling tissues or vials. All tissues were collected and frozen within approximately 5 minutes of the animal's death. The liver sections and kidneys were frozen within approximately 3-5 minutes of the animal's death. The time of euthanasia, an interim time point at freezing of liver sections and kidneys, and time at completion of necropsy were recorded. Tissues were stored at approximately -80°C or preserved in 10% neutral buffered formalin.

Tissue Collection and Processing

Liver

- 1. Right medial lobe snap frozen in liquid nitrogen and stored at ~-80°C.
- 2. Left medial lobe Preserved in 10% neutral-buffered formalin (NBF) and evaluated for gross and microscopic pathology.
- 3. Left lateral lobe snap frozen in liquid nitrogen and stored at ~-80°C.

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Heart

A sagittal cross-section containing portions of the two atria and of the two ventricles was preserved in 10% NBF. The remaining heart was frozen in liquid nitrogen and stored at \sim -80°C.

5 Kidneys (both)

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1. Left – Hemi-dissected; half was preserved in 10% NBF and the remaining half was frozen in liquid nitrogen and stored at \sim -80°C.

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2. Right – Hemi-dissected; half was preserved in 10% NBF and the remaining half was frozen in liquid nitrogen and stored at \sim -80°C.

Testes (both)

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A sagittal cross-section of each testis was preserved in 10% NBF. The remaining testes were frozen together in liquid nitrogen and stored at ~-80°C.

Brain (whole)

A cross-section of the cerebral hemispheres and of the diencephalon was preserved in 10% NBF, and the rest of the brain was frozen in liquid nitrogen and stored at \sim -80°C.

Microarray sample preparation was conducted with minor modifications, following the protocols set forth in the Affymetrix GeneChip Expression Analysis

Manual. Frozen tissue was ground to a powder using a Spex Certiprep 6800 Freezer Mill. Total RNA was extracted with Trizol (GibcoBRL) utilizing the manufacturer's protocol. The total RNA yield for each sample was 200-500 μg per 300 mg tissue weight. mRNA was isolated using the Oligotex mRNA Midi kit (Qiagen) followed by ethanol precipitation. Double stranded cDNA was generated from mRNA using the SuperScript Choice system (GibcoBRL). First strand cDNA synthesis was primed with a T7-(dT24) oligonucleotide. The cDNA was phenol-chloroform extracted and ethanol precipitated to a final concentration of 1 μg/ml. From 2 μg of cDNA, cRNA was synthesized using Ambion's T7 MegaScript in vitro Transcription Kit.

To biotin label the cRNA, nucleotides Bio-11-CTP and Bio-16-UTP (Enzo

Diagnostics) were added to the reaction. Following a 37°C incubation for six hours, impurities were removed from the labeled cRNA following the RNeasy Mini kit protocol (Qiagen). cRNA was fragmented (fragmentation buffer consisting of 200 mM Trisacetate, pH 8.1, 500 mM KOAc, 150 mM MgOAc) for thirty-five minutes at 94°C.

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Following the Affymetrix protocol, 55 µg of fragmented cRNA was hybridized on the Affymetrix rat array set for twenty-four hours at 60 rpm in a 45°C hybridization oven. The chips were washed and stained with Streptavidin Phycoerythrin (SAPE) (Molecular Probes) in Affymetrix fluidics stations. To amplify staining, SAPE solution was added twice with an anti-streptavidin biotinylated antibody (Vector Laboratories) staining step in between. Hybridization to the probe arrays was detected by fluorometric scanning (Hewlett Packard Gene Array Scanner). Data was analyzed using Affymetrix GeneChip® version 2.0 and Expression Data Mining (EDMT) software (version 1.0), GeneExpress2000, and S-Plus.

Tables 1 and 2 disclose those genes that are differentially expressed upon exposure to the named toxins and their corresponding GenBank Accession and Sequence Identification numbers, the identities of the metabolic pathways in which the genes function, the gene names if known, and the unigene cluster titles. The model code represents the various toxicity state that each gene is able to discriminate as well as the individual toxin type associated with each gene. The codes are defined in Table 4. The GLGC ID is the internal Gene Logic identification number.

Table 3 discloses those genes that are the human homologues of those genes in Tables 1 and 2 that are differentially expressed upon exposure to the named toxins. The corresponding GenBank Accession and Sequence Identification numbers, the gene names if known, and the unigene cluster titles of the human homologues are listed.

Table 4 defines the comparison codes used in Tables 1, 2, 3, and 5.

Tables 5-5CC disclose the summary statistics for each of the comparisons performed. Each of these tables contains a set of predictive genes and creates a model for predicting the renal toxicity of an unknown, *i.e.*, untested compound. Each gene is identified by its Gene Logic identification number and can be cross-referenced to a gene name and representative SEQ ID NO. in Tables 1 and 2. For each comparison of gene expression levels between samples in the toxicity group (samples affected by exposure to a specific toxin) and samples in the non-toxicity group (samples not affected by exposure

to that same specific toxin), the tox mean (for toxicity group samples) is the mean signal intensity, as normalized for the various chip parameters that are being assayed. The non-tox mean represents the mean signal intensity, as normalized for the various chip parameters that are being assayed, in samples from animals other than those treated with the high dose of the specific toxin. These animals were treated with a low dose of the specific toxin, or with vehicle alone, or with a different toxin. Samples in the toxicity groups were obtained from animals sacrificed at the timepoint(s) indicated in the Table 5 headings, while samples in the non-toxicity groups were obtained from animals sacrificed at all time points in the experiments. For individual genes, an increase in the tox mean compared to the non-tox mean indicates up-regulation upon exposure to a toxin. Conversely, a decrease in the tox mean compared to the non-tox mean indicates down-regulation.

The mean values are derived from Average Difference (AveDiff) values for a particular gene, averaged across the corresponding samples. Each individual Average Difference value is calculated by integrating the intensity information from multiple probe pairs that are tiled for a particular fragment. The normalization multiplies each expression intensity for a given experiment (chip) by a global scaling factor. The intent of this normalization is to make comparisons of individual genes between chips possible. The scaling factor is calculated as follows:

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- 1. From all the unnormalized expression values in the experiment, delete the largest 2% and smallest 2% of the values. That is, if the experiment yields 10,000 expression values, order the values and delete the smallest 200 and largest 200.
- 2. Compute the trimmed mean, which is equal to the mean of the remaining values.
- 3. Compute the scale factor SF = 100/(trimmed mean)

The value of 100 used here is the standard target valued used. Some AveDiff values may be negative due to the general noise involved in nucleic acid hybridization experiments. Although many conclusions can be made corresponding to a negative value on the GeneChip platform, it is difficult to assess the meaning behind the negative value for individual fragments. Our observations show that, although negative values are observed at times within the predictive gene set, these values reflect a real biological

phenomenon that is highly reproducible across all the samples from which the measurement was taken. For this reason, those genes that exhibit a negative value are included in the predictive set. It should be noted that other platforms of gene expression measurement may be able to resolve the negative numbers for the corresponding genes.

5 The predictive ability of each of those genes should extend across platforms, however. Each mean value is accompanied by the standard deviation for the mean. The linear discriminant analysis score (discriminant score), as disclosed in the tables, measures the ability of each gene to predict whether or not a sample is toxic. The discriminant score is calculated by the following steps:

10 Calculation of a discriminant score

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Let X_i represent the AveDiff values for a given gene across the non-tox samples, i=1...n. Let Y_i represent the AveDiff values for a given gene across the tox samples, i=1...t.

The calculations proceed as follows:

- 1. Calculate mean and standard deviation for X_i's and Y_i's, and denote these by m_x, m_y, S_X, S_Y .
- 2. For all X_i's and Y_i's, evaluate the function $f(z) = ((1/s_y)*exp(-.5*((z-m_y)/s_y)^2)) / (1/s_y)*exp(-.5*((z-m_y)/s_y)^2))$ $(((1/s_y)*exp(-.5*((z-m_y)/s_y)^2))+((1/s_x)*exp(-.5*((z-m_x)/s_y)^2))).$
- 3. The number of correct predictions, say P, is then the number of Y_i 's such that $f(Y_i) > .5$ plus the number of X_i 's such that $f(X_i) < .5$.
- 20 4. The discriminant score is then P/(n+t).

Linear discriminant analysis uses both the individual measurements of each gene and the calculated measurements of all combinations of genes to classify samples. For each gene a weight is derived from the mean and standard deviation of the toxic and nontox groups. Every gene is multiplied by a weight and the sum of these values results in a collective discriminate score. This discriminant score is then compared against collective centroids of the tox and nontox groups. These centroids are the average of all tox and nontox samples respectively. Therefore, each gene contributes to the overall prediction. This contribution is dependent on weights that are large positive or negative numbers if the relative distances between the tox and nontox samples for that gene are large and small numbers if the relative distances are small. The discriminant score for each unknown sample and centroid values can be used to calculate a probability between zero and one as to the group in which the unknown sample belongs.

Example 2: General Toxicity Modeling

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Samples were selected for grouping into tox-responding and non-tox-responding groups by examining each study individually with Principal Components Analysis (PCA) to determine which treatments had an observable response. Only groups where confidence of their tox-responding and non-tox-responding status was established were included in building a general tox model (Table 5).

Linear discriminant models were generated to describe toxic and non-toxic samples. The top discriminant genes and/or EST's were used to determine toxicity by calculating each gene's contribution with homo and heteroscedastic treatment of variance and inclusion or exclusion of mutual information between genes. Prediction of samples within the database exceeded 80% true positives with a false positive rate of less than 5%. It was determined that combinations of genes and/or EST's generally provided a better predictive ability than individual genes and that the more genes and/or EST used the better predictive ability. Although the preferred embodiment includes fifty or more genes, many pairings or greater combinations of genes and/or EST can work better than individual genes. All combinations of two or more genes from the selected list (Table 5) could be used to predict toxicity. These combinations could be selected by pairing in an agglomerate, divisive, or random approach. Further, as yet undetermined genes and/or EST's could be combined with individual or combination of genes and/or EST's described here to increase predictive ability. However, the genes and/or EST's described here would contribute most of the predictive ability of any such undetermined combinations.

Other variations on the above method can provide adequate predictive ability. These include selective inclusion of components via agglomerate, divisive, or random approaches or extraction of loading and combining them in agglomerate, divisive, or random approaches. Also the use of composite variables in logistic regression to determine classification of samples can also be accomplished with linear discriminate analysis, neural or Bayesian networks, or other forms of regression and classification based on categorical or continual dependent and independent variables.

30 Example 3: Modeling Methods

The above modeling methods provide broad approaches of combining the expression of genes to predict sample toxicity. One could also provide no weight in a

simple voting method or determine weights in a supervised or unsupervised method using agglomerate, divisive, or random approaches. All or selected combinations of genes may be combined in ordered, agglomerate, or divisive, supervised or unsupervised clustering algorithms with unknown samples for classification. Any form of correlation matrix may also be used to classify unknown samples. The spread of the group distribution and discriminate score alone provide enough information to enable a skilled person to generate all of the above types of models with accuracy that can exceed discriminate ability of individual genes. Some examples of methods that could be used individually or in combination after transformation of data types include but are not limited to: Discriminant Analysis, Multiple Discriminant Analysis, logistic regression, multiple regression analysis, linear regression analysis, conjoint analysis, canonical correlation, hierarchical cluster analysis, k-means cluster analysis, self-organizing maps, multidimensional scaling, structural equation modeling, support vector machine determined boundaries, factor analysis, neural networks, bayesian classifications, and resampling methods.

Example 4: Grouping of Individual compound and Pathology Classes

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Samples were grouped into individual pathology classes based on known toxicological responses and observed clinical chemical and pathology measurements or into early and late phases of observable toxicity within a compound (Tables 5A-5CC). The top 10, 25, 50, 100 genes based on individual discriminate scores were used in a model to ensure that combination of genes provided a better prediction than individual genes. As described above, all combinations of two or more genes from this list could potentially provide better prediction than individual genes when selected in any order or by ordered, agglomerate, divisive, or random approaches. In addition, combining these genes with other genes could provide better predictive ability, but most of this predictive ability would come from the genes listed herein.

Samples may be considered toxic if they score positive in any pathological or individual compound class represented here or in any modeling method mentioned under general toxicology models based on combination of individual time and dose grouping of individual toxic compounds obtainable from the data. The pathological groupings and early and late phase models are preferred examples of all obtainable combinations of sample time and dose points. Most logical groupings with one or more genes and one or

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more sample dose and time points should produce better predictions of general toxicity, pathological specific toxicity, or similarity to known toxicant than individual genes.

Although the present invention has been described in detail with reference to examples above, it is understood that various modifications can be made without departing from the spirit of the invention. Accordingly, the invention is limited only by the following claims. All cited patents, patent applications and publications referred to in this application are herein incorporated by reference in their entirety.

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TABLE 18 8	BUMMARÝ .				Atty: Docket No. 44921-5039W0 Doc. No. 1793397.1
Sequence ID No.	ldentifier	Conbook Ace/ Ref. Seq ID	Model Code	Gene Namo	Unigene Gluster Title
1	6949	AA012785	q		ESTs
2	25098	AA108277	h,v	 	
	-				ESTs, Highly similar to includes exons
3	17312	AA108308	lr		3 through 12 [M.musculus]
					ESTs, Moderately similar to NADH-
4	16882	AA684537	0		ubiquinone oxidoreductase subunit CI- SGDH [H.sapiens]
					ESTs, Highly similar to alpha
5	6049	AA685178	У		NAC/1.9.2. protein [M.musculus]
6	4426	AA685974	l,m		ESTs
					ESTs, Weakly similar to T23657
_	04045		_ =		hypothetical protein M01F1.6 - Caenorhabditis elegans [C.elegans]
/	21815	AA686423	9	DNA damaga indusible	Caeriornabulus elegans [C.elegans]
	4000	A A C D C 4 7 0		DNA-damage inducible transcript 3	DNA damaga indusible transgript 3
8	1600	AA686470	 	DNA-damage inducible	DNA-damage inducible transcript 3
١,	4500	A A C O C 4 7 O	l <u>.</u>	transcript 3	DNA-damage inducible transcript 3
8	1599 21997	AA686470	u	transcript 3	ESTs
9	21997	AA799325	ļ <u>u</u>		ESTs, Highly similar to AF132951 1
10	18396	AA799330	l _v		CGI-17 protein [H.sapiens]
10	19390	AA799330	 ` 		ESTs, Weakly similar to ESR1 RAT
					ESTROGEN RECEPTOR
11	6581	AA799412	f,1	<u> </u>	[R.norvegicus]
	0301	ANT 33412	1','		ESTs, Weakly similar to nucleosome
12	16538	AA799449	k		assembly protein [R.norvegicus]
12	10000	AA1 33443	 		ESTs, Moderately similar to CGI-116
13	23294	AA799472	lu		protein [H.sapiens]
14	18290	AA799497	r		ESTs
14	10230	7A1 33431			ESTs, Moderately similar to hnRNP
15	18981	AA799523	le		protein [R.norvegicus]
	20843	AA799545	h		ESTs, Weakly similar to TCPA RAT T- COMPLEX PROTEIN 1, ALPHA SUBUNIT [R.norvegicus]
16	16993	AA799560	b		ESTs
18	16576	AA799570	d		ESTs
F	100.0	7.0.00.0			ESTs, Highly similar to TBB1 RAT
			1		TUBULIN BETA CHAIN
19	18361	AA799591	li		[R.norvegicus]
20	17712	AA799598	z	***************************************	ESTs
22	18346	AA799718	f		ESTs
23	8768	AA799726	Ī		ESTs
					ESTs, Highly similar to Dgcr6 protein
24	11687	AA799732	w		[M.musculus]
25	18349	AA799744	u		ESTs
26	17494	AA799751	n		ESTs
27	18360	AA799771	General		ESTs
28	18880	AA799801	w		ESTs
					ESTs, Weakly similar to serine
29	20998	AA799803	Z		protease [R.norvegicus]
					ESTs, Highly similar to IRF7 MOUSE
					INTERFERON REGULATORY
30	21006	AA799861	С		FACTOR 7 [M.musculus]
			1	·	ESTs, Highly similar to DDRT helix-
					destabilizing protein - rat
31	15011	AA799893	General		[R.norvegicus]
					ESTs, Highly similar to 60S
امم	00044	A A 700000	_		RIBOSOMAL PROTEIN L18A
32	20811	AA799899	а		[R.norvegicus]
					ESTs, Weakly similar to S52675
					probable membrane protein YDR109c yeast (Saccharomyces cerevisiae)
22	22202	A A 700074	Gonoral		, , ,
33	23202	AA799971	General		[S.cerevisiae] ESTs, Highly similar to glycogen
24	4832	AA800190	h		phosphorylase [R.norvegicus]
34 35	21656	AA800190 AA800202	d	-	ESTs [K.norvegicus]
JJ	[21000		14	I	

TABLE 1: 8	SUMMARY *		-,		
					Dog. No. 1793997.1
ID No: Sequence	ldeniiie	ConBenk Acel Ref. Seq ID		Cene Name	Unigens Cluster IIIIs
					ESTs, Weakly similar to T15476 hypothetical protein C09F5.2 -
36	18433	AA800218	j,y,z		Caenorhabditis elegans [C.elegans]
37	6386	AA800235	u		ESTs
38	18442	AA800258	h,k		ESTs
39	21092	AA800380	у		ESTs, Weakly similar to CORTICOSTEROID 11-BETA- DEHYDROGENASE, ISOZYME 1 [R.norvegicus] ESTs, Weakly similar to glutathione
40	17325	AA800587	General cc,		peroxidase [R.norvegicus] Rattus norvegicus gene for TIS11,
41	13930	AA800613	General		complete cds
42	21372	AA800693	v v		ESTs
42	21373	AA800693	s		ESTs
43	18161	AA800701	k		ESTs
44	6595	AA800753	w		ESTs
45	13348	AA800928	General		ESTs ESTs, Highly similar to H2A1 RAT
16	23115	AA801165	lo v		HISTONE H2A.1 [R.norvegicus]
46 47	12399	AA801307	o,y General		ESTs [R:norvegicus]
48	7543	AA801395	General	· · · · · · · · · · · · · · · · · · ·	ESTs
49	24237	AA817726	t,General		ESTs
				·	ESTs, Moderately similar to T25763 hypothetical protein F46F11.4 -
50	11215	AA817921	0		Caenorhabditis elegans [C.elegans] ESTs
51	5985	AA818005	g		ESTs, Highly similar to rabkinesin-6
52	11338	AA818016	x		[M.musculus] ESTs, Weakly similar to PRSC
53	2845	AA818026	k,General		MOUSE 26S PROTEASOME REGULATORY SUBUNIT S12 [M.musculus]
54	16756	AA818089	i,k, General		ESTs, Highly similar to glycyl-tRNA synthetase [H.sapiens]
55	17771	AA818224	e,g,p, General		ESTs, Highly similar to TBB1 RAT TUBULIN BETA CHAIN [R.norvegicus]
					ESTs, Moderately similar to
56	6522	AA818261	g,m		autoantigen p542 [H.sapiens] ESTs
57 58	5924 7806	AA818359 AA818421	b.aa		ESTs ESTs
59	8237	AA818512	V		ESTs
60	17434	AA818574	h		ESTs
61	8728	AA818615	General		ESTs
62	6054	AA818658	b,v,cc, General	Diphtheria toxin receptor (heparin binding epidermal growth factor - like growth factor)	Diphtheria toxin receptor (heparin binding epidermal growth factor - like growth factor) ESTs, Moderately similar to S65785
63	11590	AA818721	d		mel-13a protein - mouse [M.musculus]
64	4291	AA818741	q,General		ESTs
65	4330	AA818747	o,General		ESTs
66	19723	AA818761	v.General		ESTs
67	13684	AA818770	h,j,l,m		Rattus norvegicus serine protease gene, complete cds
68	6322	AA818801	k		ESTs
69	7690	AA818875	General	uroguanylin	uroguanylin
70	4952	AA818907	q,General		ESTs
71	6094	AA818911	t		ESTs
	1	1	 		ESTs, Weakly similar to HP33
72	10985	AA818998	o,General		[R.norvegicus]
73	6120	AA819008	t		ESTs

Table1: S					Ally, Docket No. 44921-5099W0 Doc. No. 1798897.
Sequence ID No.	limilier	ConBenk Acel Ref. Seq (D)	Modal Gode	Gene Name	Unitgana Cluster Villa
	2500				ESTs, Weakly similar to testis specific
74	2586	AA819081	0		protein [R.norvegicus] ESTs
76 77	6438 24721	AA819269 AA819306	d,w		ESTs
	24121	AA019300	u,w		Rattus norvegicus mRNA for inositol hexakisphosphate kinase, complete
78	6250	AA819376	o,y		cds ESTs, Weakly similar to JC5707
80	6281	AA819517	j		HYA22 protein [H.sapiens]
81	10141	AA819526	<u>j</u>		ESTs
82	6551	AA819558	t		ESTs
00	6700	A A 04 00 50	_		ESTs, Moderately similar to dJ30M3.
83 84	6723 14958	AA819653 AA819744	laa		[H.sapiens]
					ESTs, Weakly similar to HS9B RAT HEAT SHOCK PROTEIN HSP 90-
85	19433	AA819776	V		BETA [R.norvegicus]
86	6204	AA819889	aa		ESTs Weakly similar to guanosine
87	22820	AA848315	General	HMm:inosine 5'-phosphate dehydrogenase 2	monophosphate reductase [R.norvegicus]
					ESTs, Weakly similar to T26686 hypothetical protein Y38F1A.6 -
88	6614	AA848389	bb		Caenorhabditis elegans [C.elegans]
89	21125	AA848437	General		ESTs
					ESTs, Moderately similar to IF4B_HUMAN EUKARYOTIC TRANSLATION INITIATION FACTOR
90	23504	AA848496	q		4B [H.sapiens] ESTs, Weakly similar to FMO1 RAT
91	18532	AA848675	g		DIMETHYLANILINE MONOOXYGENASE [R.norvegicus]
92	21140	AA848738	c	1	ESTs
					ESTs, Moderately similar to AF13294
93	16128 22923	AA848807	0		1 CGI-12 protein [H.sapiens] ESTs
94 95	17339	AA848929 AA849497	g General		ESTS
96	11727	AA849518	General		ESTs
30	111727	74043510	i,I,m,		1013
97	21275	AA849796	General		ESTs
98	16678	AA849827	aa		ESTs
99	8515	AA849917	е		ESTs
100	18447	AA849939	General		ESTs
101	12130	AA850037	P	<u> </u>	ESTs
102	23981	AA850040	x,aa	cyclase-associated protein homologue	cyclase-associated protein homologue
103	13615	AA850364	t		MOUSE RAS-RELATED PROTEIN RAB-17 [M.musculus]
105	2637	AA850893	×		ESTs, Highly similar to hypothetical protein [H.sapiens]
106	22093	AA850909	lâ		ESTs
107	21766	AA850916	С		ESTs
108	2847	AA850919	w		ESTs, Weakly similar to dithiolethione inducible gene-1 [R.norvegicus] Rattus norvegicus mRNA for ras-
					GTPase-activating protein SH3-
109	12162	AA850975	h		domain binding protein, partial cds
110	9514	AA850978	General		ESTs
111	3924	AA851017	e,q		ESTs, Highly similar to molybdopterin synthase large subunit [M.musculus]
111	3925	AA851017	o,General		ESTs, Highly similar to molybdopterin synthase large subunit [M.musculus]
112	4490	AA851184	a,k		Rattus norvegicus mRNA for cathepsin Y, partial cds

TABLE 1: S					Ally. Docket No. 44921-5039W0
	Hilly Still - I was				Doc. No. 1793997.1
Sequence ID No. :	ldeniiier	GenBank Acc/ Ref. Seq ID	Model Gode	©ene Name	Unigene Cluster Title
		MILANDESOCKA MILANDESOCIA (MILANDESOCIA (MILANDESOCIA) (MILANDESOCIA (MI	The second secon		ESTs, Weakly similar to T28050
					hypothetical protein ZK856.11 -
113	19187	AA851230	General		Caenorhabditis elegans [C.elegans]
					ESTs, Highly similar to ubiquitin
	10100	4 4 0 5 4 0 0 7	_		specific protease UBP43
114	19189	AA851237	С		[M.musculus] ESTs, Highly similar to hypothetical
115	15386	AA851241	m		protein [H.sapiens]
110	10000	70.001211			ESTs, Weakly similar to A61382
			g,I,		phosphorylation regulatory protein HP-
116	21462	AA851261	General		10 [H.sapiens]
117	21471	AA851343	General		ESTs
				HHs:NADH dehydrogenase	ESTs, Moderately similar to
				(ubiquinone) Fe-S protein 8	NUIM_HUMAN NADH-UBIQUINONE
140	40000	4 4 0 5 4 0 7 0	<u> </u>	(23kD) (NADH-coenzyme Q	OXIDOREDUCTASE 23 KD SUBUNIT
118	16902	AA851379	P	reductase)	PRECURSOR [H.sapiens] ESTs, Moderately similar to kinesin-
119	23376	AA851392	i,x		like DNA binding protein [H.sapiens]
110	20070	7 0 00 1002	···^	 	ESTs, Moderately similar to kinesin-
119	23377	AA851392	l _x		like DNA binding protein [H.sapiens]
120	13349	AA851417	General	****	ESTs
121	21527	AA851733	r,u		ESTs
			i,o,u,		Rattus norvegicus osteoactivin mRNA
122	4048	AA851814	General		complete cds
					ESTs, Highly similar to SSRA HUMAN
	1				TRANSLOCON-ASSOCIATED
400	40564	AA054074	bb		PROTEIN, ALPHA SUBUNIT PRECURSOR [H.sapiens]
123	10561	AA851871	UD		Rattus norvegicus CaM-kinase II
124	17411	AA858621	l j,y		inhibitor alpha mRNA, complete cds
127	1771	701030021	V.7		ESTs, Weakly similar to MCM6 RAT
					DNA REPLICATION LICENSING
125	1801	AA858636	k,s,x,bb		FACTOR MCM6 [R.norvegicus]
126	18350	AA858674	р		ESTs
127	19484	AA858693	е		ESTs
128	6360	AA858696	d		ESTs
400	47204	4 4 0 5 0 7 0 4	<u> </u>		ESTs, Weakly similar to Reg receptor
129	17334	AA858704	p		[R.norvegicus] ESTs, Weakly similar to dJ413H6.1.1
130	6380	AA858758	l _a		[H.sapiens]
131	13219	AA858759	a		ESTs
	10210	7 0 10007 00	I.m.Gener		
132	6384	AA858788	al		ESTs
	_				ESTs, Highly similar to p40 seven-
					transmembrane-domain protein
134	13412	AA858830	p		[M.musculus]
135	7279	AA858892	ļ		ESTs
136	18217	AA858930	t		ESTS TARRACINIVI TRNA
			1	HHe:asparagicul-tDNA	ASPARAGINYL-TRNA SYNTHETASE, CYTOPLASMIC
137	5867	AA858953	v General	HHs:asparaginyl-tRNA synthetase	[H.sapiens]
101	10007	, 0.000300	7,001161a1	7.1.101.000	ESTs, Moderately similar to 156526
			1		interleukin 1 receptor type I - rat
138	14479	AA858969	r		[R.norvegicus]
139	6431	AA859085	t		ESTs
140	17361	AA859114	o,General	<u> </u>	ESTs
141	21025	AA859241	General	outer membrane protein	outer membrane protein
142	10076	AA859271	С		ESTs Weakly similar to CYSR RAT
					CYSTEINE-RICH PROTEIN 1
143	21791	AA859333	k		[R.norvegicus]
1-70	-1/3	, , , , , , , , , , , , , , , , , , , ,	cc,Gener		[
144	16314	AA859348	al		ESTs
145	18862	AA859520	f		ESTs
146	15059	AA859545	r		ESTs

TABLE 1: S	WININARY:				Ally. Docket No. 44921-5039W0 Doc. No. 1793897.1
Sequence ID No.	ldeniiier	CenBank Acc	Model Code	Gene Namo	Unigase Civeter Tille
			1		Rattus norvegicus late gestation lung
147	19894	AA859581	s		protein 1 (Lgl1) mRNA, complete cds
148	14353	AA859585	h		ESTs
140	16210	AA859648	h		ESTs, Weakly similar to DnaJ homolog 2 [R.norvegicus]
149 150	16318 17316	AA859652	General		ESTs
151	19067	AA859663	n,q		ESTs
152	22406	AA859680	n		ESTs
153	20599	AA859690	x		ESTs
454	14004				ESTs, Weakly similar to YNH2_CAEEL HYPOTHETICAL 31.0 KD PROTEIN R107.2 IN
154	14261	AA859693	lu 	HHs:protoporphyrinogen	CHROMOSOME III [C.elegans] ESTs, Highly similar to PPOX MOUSE PROTOPORPHYRINOGEN OXIDASE
155	14138	AA859700	v	oxidase	[M.musculus]
				HHs:protoporphyrinogen	ESTs, Highly similar to PPOX MOUSE PROTOPORPHYRINOGEN OXIDASE
155	14139	AA859700	V	oxidase	[M.musculus] ESTs, Weakly similar to IF4E MOUSE EUKARYOTIC TRANSLATION INITIATION FACTOR 4E
157	22374	AA859804	1		[R.norvegicus] ESTs, Moderately similar to LYOX
			l		RAT PROTEIN-LYSINE 6-OXIDASE
158	22385	AA859805	b,k		PRECURSOR [R.norvegicus]
159 160	22773 22816	AA859885 AA859898	n k,x,z		ESTs ESTs
161	11891	AA859926	X		ESTs
 	11001	741000020	<u> </u>		ESTs, Highly similar to N-
162	23070	AA859942	k		myristoyltransferase 1 [M.musculus]
163	23121	AA859948	k		ESTs
164	23166	AA859954	cc,Gener al		ESTs
165	18468	AA859966	aa		ESTs, Weakly similar to Edp1 protein [M.musculus]
166	23336	AA859981	q	HHs:inositol(myo)-1(or 4)- monophosphatase 2	MYO-INOSITOL-1(OR 4)- MONOPHOSPHATASE [R.norvegicus]
167	4222	AA860024	a,bb_		ESTs, Highly similar to EF1G_HUMAN ELONGATION FACTOR 1-GAMMA [H.sapiens]
168	13974	AA860030	u,x,Gener al		Rattus norvegicus mRNA for class I beta-tubulin, complete cds
169	7090	AA860039	x	receptor (RHAMM)	EST, Hyaluronan mediated motility receptor (RHAMM) ESTs, Moderately similar to T08661
170 171	23769	AA860055	k,x w		anti-silencing protein ASF1 homolog DKFZp547E2110.1 [H.sapiens]
171	16323	AA866240	W		ESTs, Weakly similar to PE2R RAT 20 ALPHA-HYDROXYSTEROID
172	4462	AA866264	General		DEHYDROGENASE [R.norvegicus] ESTs, Weakly similar to A60543
173	15884	AA866276	k	4-hydroxyphenylpyruvic acid	protein kinase [R.norvegicus] 4-hydroxyphenylpyruvic acid
174	17742	AA866302	с,у	dioxygenase Solute carrier family 4,	dioxygenase Solute carrier family 4, member 1,
175	16333	AA866414	a,h	member 1, anion exchange protein 1 (kidney band 3)	anion exchange protein 1 (kidney band 3) ESTs, Moderately similar to AF141884
176	18918	AA866444	p,q		1 oligophrenin-1 like protein [H.sapiens]
177	16853	AA866454	j,l,m,y,z		ESTs
178	18995	AA866459	h,m_	L	ESTs

TABLE 1: St					Atty: Docket No. 44921-5039000 Doc. No. 1793397.1
Sequence ID No.	ldentifier	Cenbank Acc Ref. Seq ID	Model Code	Gene Name	Unigane Gluster Tille
					ESTs, Highly similar to FGD1 MOUSE PUTATIVE RHO/RAC GUANINE NUCLEOTIDE EXCHANGE FACTOR
179	16013	AA866482	s		[M.musculus]
180	26036	AA874849	r		
181	16059	AA874857	h		ESTs
182	16069	AA874873	r		ESTs
400					ESTs, Weakly similar to RNA binding
183 184	21633 16192	AA874951 AA874995	w		protein [H.sapiens] ESTs
			<u></u>		ESTs, Highly similar to RET3 BOVIN RETINOIC ACID-BINDING PROTEIN
185	16254	AA875025	i		I, CELLULAR [R.norvegicus]
100	16313	A A 975020	cc,Gener		ESTs
186 187	16312 20701	AA875032 AA875097	al b		Rat alpha-fibrinogen mRNA, 3' end
					ESTs, Highly similar to ARF3_HUMAN ADP-RIBOSYLATION FACTOR
188	16416	AA875098	bb		[R.norvegicus] ESTs, Highly similar to RUXE_HUMAN SMALL NUCLEAR RIBONUCLEOPROTEIN E
189	16419	AA875102	bb		[M.musculus]
			l,m,		
190	15313	AA875126	General		ESTs
					ESTs, Weakly similar to AF151834 1
191 192	10936 18084	AA875146 AA875186	h h		CGI-76 protein [H.sapiens]
192	10004	AA073100	11		ESTS, Highly similar to IF39_HUMAN EUKARYOTIC TRANSLATION INITIATION FACTOR 3 SUBUNIT 9
193	15371	AA875205	u		[H.sapiens]
194	15401	AA875257	x,z		ESTs
				HHs:NADH dehydrogenase (ubiquinone) Fe-S protein 7 (20kD) (NADH-coenzyme Q	ESTs, Highly similar to NUKM
195	15410	AA875268	p,s	reductase)	HUMAN, partial CDS [H.sapiens]
196	15420	AA875286	f		ESTs
197	15446	AA875327	s,w		ESTs
198	7936	AA875495	b,General		ESTs ESTs, Highly similar to includes exons
199	17314	AA875509	i,I,m		3 through 12 [M.musculus] ESTs, Highly similar to MLES RAT
	0.4.70				MYOSIN LIGHT CHAIN ALKALI, SMOOTH-MUSCLE ISOFORM
200	24472	AA875523	k		[R.norvegicus] ESTs
202	15587 15617	AA875577 AA875620	J General		ESTs
202	15618	AA875620	General		ESTs
203	5384	AA891041	f,cc,Gene ral	jun B proto-oncogene	jun B proto-oncogene
204	24814	AA891209	f,p		ESTs, Moderately similar to R33729 1, partial CDS [H.sapiens] ESTs, Weakly similar to AF151373 1
205	21930	AA891322	d		nucleolin-related protein NRP [R.norvegicus]
206	17225	AA891553	h		ESTs, Highly similar to eIF3 p66 [M.musculus]
					ESTs, Weakly similar to S67314 regulatory protein RMS1 - yeast (Saccharomyces cerevisiae)
207	7522	AA891571	j,m		[S.cerevisiae]
208	9071	AA891578	b		ESTs
209	19321	AA891666	u	melanoma antigen, family D, 1	melanoma antigen, family D, 1

	SUMMARY				Auy. Docket No. 44921-5039000 Doc. No. 1793397.1
Sequence ID No.	ldenililer	Conbank Acci Ref. Seq (D)	Model Code	Cone Name	Unitene Cluster Title
210	17693	AA891737	j,l,m,n,y,z		ESTs
211	17256	AA891739	General		ESTs, Weakly similar to T22521 hypothetical protein F52H3.5 - Caenorhabditis elegans [C.elegans] ESTs, Moderately similar to FINC RAT
213	18269	AA891769	General		FIBRONECTIN PRECURSOR [R.norvegicus]
214	9905	AA891774	s,bb, D239Gen eral		ESTs
215	17061	AA891812	d		ESTs, Highly similar to alpha-adducin, hypertensive phenotype [R.norvegicus]
216	7050	AA891824	h		Rattus norvegicus clone ZG52 mRNA sequence
217	4463	AA891831	General		ESTs, Weakly similar to PE2R RAT 20 ALPHA-HYDROXYSTEROID DEHYDROGENASE [R.norvegicus]
218 219	14289	AA891838	i		ESTs, Highly similar to muscle protein 684 [M.musculus] ESTs
	20523	AA891842	r,cc		ESTS, Moderately similar to ACY1_HUMAN AMINOACYLASE-1 [H.sapiens]
220	17779 17438	AA891914 AA891943	g,s,z General		ESTs
221	22862	AA891944			ESTs
222 223	1159	AA891949	e,z		ESTs
224	4473	AA891965	General		ESTs, Weakly similar to T31496 hypothetical protein Y116A8C.25 - Caenorhabditis elegans [C.elegans] ESTs, Highly similar to chromatin
225	6362	AA892053	f,j,l,m	1711	structural protein homolog Supt5hp [M.musculus]
226	9037	AA892066	y General		ESTs ESTs, Weakly similar to proline dehydrogenase [M.musculus]
227 228	19469 14595	AA892112 AA892128	o,t,v	· · · · · · · · · · · · · · · · · · ·	ESTs
229	16527	AA892154	cc c	,	ESTs
230	4482	AA892173	bb		EST
231	20917	AA892238	h		ESTs ESTs, Weakly similar to PC4221
232	2357	AA892268	d		protein-tyrosine kinase [R.norvegicus]
233	18183	AA892271	h	· · · · · · · · · · · · · · · · · · ·	ESTs
234	6523	AA892299	d		ESTs ESTs, Highly similar to RL3 RAT 60S
236	13647	AA892367	а		RIBOSOMAL PROTEIN L3 [R.norvegicus] ESTs, Highly similar to AF151893 1
237	3473	AA892378	v		CGI-135 protein [H.sapiens]
238	17682	AA892382	j,p,s,x, General		ESTs, Moderately similar to AF185570 1 putative N-acetyltransferase Camello 4 [R.norvegicus]
239	820	AA892395	g,s	Aldolase B, fructose- biphosphate	Aldolase B, fructose-biphosphate
240	14754	AA892414	u		ESTs
241	17439	AA892446	f		ESTS ESTS, Moderately similar to UCRY_HUMAN UBIQUINOL- CYTOCHROME C REDUCTASE COMPLEX 6.4 KD PROTEIN
242	16469	AA892462	p : Canasal		[H.sapiens] Rattus norvegicus mRNA for prostasir
243	13609	AA892468	i,General n,v,Gener		Precursor, complete cds Rattus norvegicus mRNA for prostasir
243	13610	AA892468	al	<u> </u>	precursor, complete cds

Table 1: S	UMMARY				Aiiy, Docket No. 44924-5039W Doc. No. 1793397,
Sequence ID No.	lieniiier	ConBank Acci Ref. Seq ID	Modal Modal	Cone Name	Unigane Civetar Title
	0054				ESTs, Highly similar to HISTONE
244 245	9254 11991	AA892470 AA892483	n,u s		H2A.Z [R.norvegicus]
245	11991	AA692463	5		ESTs, Moderately similar to LYAG MOUSE LYSOSOMAL ALPHA- GLUCOSIDASE PRECURSOR
246	1522	AA892486	f		[M.musculus] ESTs, Moderately similar to S63540
247	11994 _	AA892507	aa		protein DS 1, 24K [H.sapiens]
248	23888	AA892520	w		ESTs
248	23889	AA892520	h		ESTs
249	8599	AA892522	р		ESTs
250	15154	AA892532	р		R.norvegicus (Wistar) CaBP1 mRNA ESTs, Highly similar to multi- membrane spanning polyspecific
251	17468	AA892545	r		transporter [M.musculus] ESTs, Highly similar to ras-GTPase- activating protein SH3-domain binding
252	11203	AA892554	f,h a,bb,		protein [M.musculus] ESTs, Moderately similar to PTD012
253	18906	AA892561	General		[H.sapiens] R.norvegicus mRNA for nucleolar
254	19327	AA892562	f,j,y,z		protein NAP57
255	18274	AA892572	р		ESTs
256	4512	AA892578	CC		ESTs ESTs, Highly similar to RL8_HUMAN 60S RIBOSOMAL PROTEIN L
257	15876	AA892582	w		[R.norvegicus]
258	19085	AA892598	General		ESTs
258	19086	AA892598	General		ESTs ESTs, Highly similar to H4 HUMAN
259	20065	AA892647	ı		HISTONE H4 [R.norvegicus]
260	20088	AA892666	a,n		ESTs
261	23783	AA892773	n		ESTs Rat mitochondrial proton/phosphate
262	17549	AA892776	f,z		symporter mRNA, complete cds
263	13542	AA892798	b	HHs:glyoxylate reductase/hydroxypyruvate	ESTs ESTs, Weakly similar to SERA RAT D 3-PHOSPHOGLYCERATE
264	22537	AA892799	General	reductase HHs:glyoxylate	DEHYDROGENASE [R.norvegicus] ESTs, Weakly similar to SERA RAT D 3-PHOSPHOGLYCERATE
264	22539	AA892799	v	reductase/hydroxypyruvate reductase HHs:glyoxylate	DEHYDROGENASE [R.norvegicus] ESTs, Weakly similar to SERA RAT D
264	22538	AA892799	General	reductase/hydroxypyruvate reductase	3-PHOSPHOGLYCERATE DEHYDROGENASE [R.norvegicus] ESTs, Weakly similar to S70642
265	6951	AA892820	h		ubiquitin ligase Nedd4 - rat [R.norvegicus]
266	23322	AA892821	j,z		Rattus norvegicus aiar mRNA for androgen-inducible aldehyde reductase, complete cds
267	17923	AA892843	f		ESTs, Weakly similar to T29904 hypothetical protein F59A3.3 - Caenorhabditis elegans [C.elegans]
268	22871	AA892859	m		ESTs, Weakly similar to procollagen- lysine 5-dioxygenase [R.norvegicus]
269	9053	AA892861	p,v, General		ESTs
					ESTs, Weakly similar to EF2 RAT ELONGATION FACTOR 2
270	16482	AA892940	w		[R.norvegicus] Rattus norvegicus HP33 mRNA,
271 272	12020	AA893035	j.y General		complete cds ESTs
	3863	AA893060	usenerai	i .	IEO18

Market in	BUMMARY				/Aiiy. Docket No. 44921-5039\\ Doc. No. 1793397.
Sequence ID No.	ldeniller	Conbank Ace Ref. Seq ID	Model Code	Gene Name	Unigene Auster Tille
274	21305	AA893082	General		ESTs
275	16591	AA893191	j,z		ESTs
276	17447	AA893192	General		ESTs
277	3876	AA893205	n		ESTs
278	3878	AA893230	General		ESTs, Weakly similar to CALM_HUMAN CALMODULIN [R.norvegicus]
279	20986	AA893242	q	Acyl CoA synthetase, long chain	Acyl CoA synthetase, long chain
	1		i,z,		ESTs, Moderately similar to
280	16168	AA893280	General		adipophilin [H.sapiens]
281	3886	AA893289	j,m,y		ESTs
282	15209	AA893327	У		ESTs
283	17800	AA893436	СС		ESTs
284 285	17836 9084	AA893626 AA893717	h x		ESTs, Weakly similar to LIS1 MOUSE PLATELET-ACTIVATING FACTOR ACETYLHYDROLASE IB ALPHA SUBUNIT [R.norvegicus]
286	22731	AA893743	d		ESTs
287	12031	AA893860	v	HHs:threonyl-tRNA synthetase	ESTs, Moderately similar to SYTC_HUMAN THREONYL-TRNA SYNTHETASE, CYTOPLASMIC [H.sapiens]
288	17897	AA893905	k	 -	ESTs
289	3447	AA893982	d		ESTs
290	22583	AA894009	n		1
291	10540	AA894027	<u> </u>		EST
292	4569	AA894059	x		ESTs, Highly similar to A55748 protein kinase [M.musculus] ESTs, Weakly similar to APP2 RAT
293 294	18419 17336	AA894130 AA894297	d		AMYLOID-LIKE PROTEIN 2 PRECURSOR [R.norvegicus] ESTs
295	19120	AA894318	f,j		ESTs
296	19762	AA899113	11,,		ESTs .
296		AA899219	lu u		Rat mRNA for beta-tubulin T beta15
298	18286	AA899498	w		ESTs, Weakly similar to T26581 hypothetical protein Y32B12A.3 - Caenorhabditis elegans [C.elegans]
298	22052	AA899498	q		ESTs, Weakly similar to T26581 hypothetical protein Y32B12A.3 - Caenorhabditis elegans [C.elegans]
299	21628	AA899563	aa		ESTs
300	4262	AA899590	ļi		ESTs
301 302	4661 21354	AA899709 AA899721	t,General	receptor activity modifying protein 3	receptor activity modifying protein 3 ESTs
			q		Rattus norvegicus epidermal growth factor receptor related protein (Errp)
303	17905	AA899762	General		mRNA, complete cds
304	15231	AA899840	r		ESTs
				topoisomerase (DNA) II	
305	23778	AA899854	c,k,x	alpha	topoisomerase (DNA) II alpha
306	22060	AA899898	b		ESTs
307	9114	AA899951	v,General		ESTs
308	8988	AA900148	f		ESTs Rattus norvegicus mRNA for
309	11841	AA900247	v_		Hsp70/Hsp90 organizing protein ESTs, Highly similar to ALPHA-2-
210	4725	AA000300	66		MACROGLOBULIN PRECURSOR
310	4725	AA900290	Coperal		[R.norvegicus]
311	4747	AA900465	General	ļ	ESTs ESTs
312	20988	AA900562	0		
313	3822	AA900863	b,g, General		ESTs, Weakly similar to nuclear RNA helicase [R.norvegicus]

TABLE 18	SIUMIWASA				A1117. Dooket No. 44921-5039000
. بر حقاظی ا	բառուհում ։				Doc No. 1793897.1
Seguence		GenBank Ace/	(Model)		
D NO -	ldeniiiler	Ref. Seq ID	ලිලුල්ල	Gene Name	Unigene Cluster Tille
Linder Comment				duning the same and the same an	ESTs, Weakly similar to T20702
					hypothetical protein F10C2.6 -
315	12420	AA901017	b		Caenorhabditis elegans [C.elegans]
316	4849	AA901155	s		Rattus norvegicus CDK105 mRNA
					ESTs, Highly similar to IF2B_HUMAN EUKARYOTIC TRANSLATION
					INITIATION FACTOR 2 BETA
317	3959	AA901338	General		SUBUNIT [H.sapiens]
					ESTs, Highly similar to ATP-specific
l			l		succinyl-CoA synthetase beta subunit
318 319	22846 4895	AA923982 AA923999	a,d k		[M.musculus] ESTs
319	4695	AA923999	cc.		2318
320	21546	AA924188	General		ESTs
					<u> </u>
321	24192	AA924210	n,General		ESTs
222	4022	1 4004204	g,I,Gener		FOT
322	4933	AA924301	al		ESTs, Moderately similar to
					NO56 HUMAN NUCLEOLAR
323	4944	AA924405	I,General		PROTEIN NOP56 [H.sapiens]
324	4948	AA924428	r		ESTs
					ESTs, Weakly similar to NPT2 RAT
					RENAL SODIUM-DEPENDENT PHOSPHATE TRANSPORT
325	4949	AA924432	General		PROTEIN 2 [R.norvegicus]
326	18891	AA924598	е		ESTs
			1	HHs:glyoxylate	ESTs, Weakly similar to SERA RAT D-
				reductase/hydroxypyruvate	3-PHOSPHOGLYCERATE
327	22540	AA924630	v,General		DEHYDROGENASE [R.norvegicus]
				HHs:glyoxylate reductase/hydroxypyruvate	ESTs, Weakly similar to SERA RAT D- 3-PHOSPHOGLYCERATE
327	22541	AA924630	General	reductase	DEHYDROGENASE [R.norvegicus]
328	14759	AA924766	k		ESTs
329	23123	AA924794	х		ESTs
330	4067	AA924813	g,p		ESTs
331	2888	AA924902	r,General		ESTs ESTs, Highly similar to sec7 domain
332	18130	AA924964	d		family member [H.sapiens]
333	23141	AA925019	r		ESTs
					ESTs, Weakly similar to MCT7 RAT
004	00405	44005000	0		MAST CELL PROTEASE 7
334	23195	AA925026	General f.aa.		PRECURSOR [R.norvegicus]
335	21458	AA925049	General		ESTs
					ESTs, Moderately similar to S20710
					hypothetical protein, 16K - mouse
336	5073	AA925061	m		[M.musculus]
337	14790	AA925087	o,General		ESTs
557	17730	7.57.5001	O, Octional		EST, Highly similar to T50621
					hypothetical protein DKFZp7620076.1
338	5089	AA925126	g		[H.sapiens]
					ESTs, Moderately similar to BHMT
	İ				RAT BETAINEHOMOCYSTEINE S- METHYLTRANSFERASE
339	23261	AA925145	k,General		[R.norvegicus]
		1	,, _ 5,,,,,,,,,		ESTs, Moderately similar to
					neurodegeneration-associated protein
340	17363	AA925150	а	ļ	1 [R.norvegicus]
341 342	23448 23159	AA925167 AA925318	l e	I-kappa-B-beta	ESTs I-kappa-B-beta
343	21500	AA925353	k	I-vahha-p-nera	ESTs
344	22479	AA925418	t		ESTs
345	21151	AA925539	b		ESTs

				-62-	
TABLE 1:	SUMMARY		644 (1) (1) (1) (1) (1) (1) (1) (1) (1) (1)		ANNY, Docket No. 44921-50:9000 Doc. No. 1793397.1
equeño D No.	e lideniiier	CinBank Ace Ref. Seq ID		Cene Name	Unigan Cluster Title
346	16944	AA925541	f	heterogeneous nuclear ribonucleoprotein L	heterogeneous nuclear ribonucleoprotein L
				heterogeneous nuclear	heterogeneous nuclear
346	16945	AA925541	<u> t</u>	ribonucleoprotein L HHs:succinate	ribonucleoprotein L ESTs, Highly similar to
347	17514	AA925554	bb	dehydrogenase complex, subunit A, flavoprotein (Fp)	DHSA_HUMAN SUCCINATE DEHYDROGENASE [H.sapiens]
348	5183	AA925662	i,General	Subulit A, havoprotein (i p)	ESTs
349	23189	AA925844	r		ESTs
				**	ESTs, Highly similar to IMB3_HUMAN IMPORTIN BETA-3 SUBUNIT
350	23190	AA925863	aa		[H.sapiens]
351	5252	AA926051	General		EST ESTs
352 353	22967 17157	AA926080 AA926129	h,cc b		ESTS
JJJ	17 137	777320123			
354	13411	AA926196	u,General	putative potassium channel	ESTs
355	5295	AA926247	General	TWIK	putative potassium channel TWIK
					ESTs, Moderately similar to NEURONAL PROTEIN 3.1
356	22928	AA926262	General		[M.musculus] ESTs, Moderately similar to T13963
357	8948	AA926316	г		formin related protein, lymphocyte specific - mouse [M.musculus]
358	21798	AA926365	aa		ESTs, Moderately similar to AF151827 1 CGI-69 protein [H.sapiens]
359	9942	AA942697	s		ESTs
360	6039	AA942716	x,General		ESTs, Highly similar to HN1 [M.musculus]
361	11174	AA942745	g,o,w		ESTs
362	23005	AA942770	g		ESTs
363	21318	AA942774	General		ESTs ESTs, Weakly similar to T26686
364	6615	AA942889	V		hypothetical protein Y38F1A.6 - Caenorhabditis elegans [C.elegans]
304	0013	741342003	· ·		ESTS, Highly similar to KFMS RAT MACROPHAGE COLONY STIMULATING FACTOR I
365	6691	AA943028	c		RECEPTOR PRECURSOR [R.norvegicus]
366	22142	AA943066	Р		ESTs, Weakly similar to p68 RNA helicase [R.norvegicus]
367	21993	AA943149	v,General		ESTs, Weakly similar to T00084 hypothetical protein KIAA0512 [H.sapiens]
368	9061	AA943508	General		ESTs, Weakly similar to T08666 hypothetical protein DKFZp547N0510.1 [H.sapiens]
369	24390	AA943531	b,j,n,y		ESTs, Weakly similar to VIL1 MOUSE VILLIN [M.musculus]
370	13976	AA943532	f,s,x		Rattus norvegicus mRNA for class I beta-tubulin, complete cds
371	22248	AA943537	cc, General		Rattus norvegicus zyxin mRNA, partia
					ESTs, Highly similar to T2DA_HUMAN TRANSCRIPTION INITIATION FACTOR TFIID 20/15 KDA
372	22257	AA943558	m u,cc,		SUBUNITS [H.sapiens]
373	12673	AA943773	General		ESTs
374	13641	AA944154	u		ESTs
375	2658				ESTs

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Sequence ID No.	ldeniller:	Conbank Acc Rof. Seg ID	Model Gode	Cene Name	Unigene Cluster Tille
					ESTs, Highly similar to CKS2 MOUSE CYCLIN-DEPENDENT KINASES REGULATORY SUBUNIT 2
377	20903	AA944180	i,x		[M.musculus]
378	13507	AA944244	v		ESTs
379	15596	AA944353	General		ESTs
380	22681	AA944413	i,v,cc, General		ESTs
	1				ESTs, Highly similar to hypothetical
381	6711	AA944439	General		protein [M.musculus] ESTs, Weakly similar to FIBA RAT
			i,q,		FIBRINOGEN ALPHA/ALPHA-E
382	14763	AA944481	General h		CHAIN PRECURSOR [R.norvegicus] ESTs
383	22466	AA944605	in .		ESTs, Weakly similar to A44437
					regenerating liver inhibitory factor
384	12301	AA944727	b	(2)(1)	RL/IF-1 - rat [R.norvegicus]
]		HHs:polymerase (RNA) II	ESTs, Highly similar to RNA polymerase II 23kD subunit
205	7023	AA944792	d.m.aa	(DNA directed) polypeptide E (25kD)	IfH.sapiensl
385 386	22536	AA944803	bb	(2380)	ESTs
387	22501	AA944811	g,l		ESTs
388	23967	AA944831	s	****	ESTs
389	26084	AA944922	i		
390	11974	AA944958	General		ESTs
391	22547	AA944970	aa		ESTs
392	22554	AA945076	z,General		ESTs
393	14352	AA945181	General		ESTs
395	1798	AA945569	General		R.norvegicus alpha-1-macroglobulin mRNA, complete cds
396	22050	AA945604	i,aa		ESTs
397	19731	AA945615	d,o		ESTs
					ESTs, Weakly similar to DHQU RAT NAD(P)H DEHYDROGENASE
398	22612	AA945624	a,General		[R.norvegicus]
399	22618	AA945656	aa		ESTs
400	11871	AA945679	Congrel		ESTs ESTs
401 402	22656 6720	AA945818 AA945828	General p		ESTs
403	22351	AA945867	m		ESTs
404	22665	AA945877	f		ESTs
405	24243	AA945950	Ь		ESTs
406	22689	AA945962	General		ESTs
407	22692	AA945986	d		ESTs
408	22696	AA945996	c,General		ESTs
408	22697	AA945996	c,o		ESTs
409	22658	AA945998	w		ESTs
				LINATURE DE LA CONTRA	ESTs, Highly similar to COXG
410	20822	AA946040	s	HMm:RIKEN cDNA 2010000G05 gene	MOUSE CYTOCHROME C OXIDASE POLYPEPTIDE VIB [M.musculus]
410 411	20832 18337	AA946046	General	20 10000000 gene	ESTs
	1.0007		Contrai		Rattus norvegicus laminin-5 alpha 3
412	825	AA946108	General		chain mRNA, complete cds
413	8639	AA946221	e,cc, General		ESTs
414	23237	AA946224	f		ESTs
415	15600	AA946250	o,aa		ESTs
416	19387	AA946275			[ESTs, Highly similar to AR21_HUMAN ARP2/3 COMPLEX 21 KD SUBUNIT [H.sapiens]
417	6351	AA946344	d	PCTAIRE-1 protein kinase, alternatively spliced	PCTAIRE-1 protein kinase, alternatively spliced
				anomialitory opilioco	ESTs, Highly similar to autoantigen
418	22057	AA946348	е		[H.sapiens]
419	22069	AA946349	aa		ESTs

TABLE 1:	SUMMARY				Ally, Docket No. 44921-5039WC Doc. No. 1793397.1
Seguence ID No.	. [dentifier	GenBank Acc/ Ref. Seq ID	Model Code	Geno Namo	Unigene Cluster Tible
420	13962	AA946351	General		ESTs
					ESTs, Highly similar to Ring3
421	18280	AA946361	g		[M.musculus]
422	18944	AA946391	V		ESTs ESTs, Moderately similar to p18
424	21410	AA946408			component of aminoacyl-tRNA synthetase complex [H.sapiens]
425	643	AA946439	o,y		Rat H4 gene for somatic histone H4
426	20736	AA946443			ESTS, Highly similar to NPD1 MOUSE NEURAL PROLIFERATION DIFFERENTIATION AND CONTROL PROTEIN-1 PRECURSOR [M.musculus]
427	21878	AA946448	X r		ESTs
721	21070	77,040440	<u> </u>		ESTs, Highly similar to AF151863 1
428	21947	AA946451	bb		CGI-105 protein [H.sapiens]
429	17499	AA946467	General		ESTs
- 		T	1		Rat mRNA for alpha-2u globulin-
430	1809	AA946503	x,General		related protein
431	23360	AA955104	f		ESTs
432	23471	AA955162	General		ESTs
433	9452	AA955206	b,General		ESTs
434	23512	AA955282	General		ESTs
435	22596	AA955298	General	lingaratain hinding protein	ESTs lipoprotein-binding protein
436 437	23283	AA955391	h General	lipoprotein-binding protein	ESTs
437	23546	AA955393	General		ESTs, Weakly similar to SX10 RAT
438	12404	AA955408	b		TRANSCRIPTION FACTOR SOX-10 [R.norvegicus]
439	23626	AA955540	aa		ESTs
441	17540	AA955914	bb		EST,EST, Moderately similar to FBRL MOUSE FIBRILLARIN [M.musculus],ESTs, Highly similar to FBRL MOUSE FIBRILLARIN [M.musculus]
442	24277	AA955962	General		ESTs
443	19939	AA955980	General		ESTs, Moderately similar to pescadillo [H.sapiens]
444	24000	AA956005	i		ESTs, Weakly similar to AF139894 1 RNA-binding protein alpha-CP1 [M.musculus]
445	11050	AA956164	s,v		ESTs, Weakly similar to TCPA RAT T- COMPLEX PROTEIN 1, ALPHA SUBUNIT [R.norvegicus]
446	498	AA956278	a,General		ESTs
447	23409	AA956294	q		ESTs
449	23773	AA956476	f,x		ESTs ESTs, Highly similar to ET putative
450	23799	AA956530	d		translation product [M.musculus]
451	23800	AA956534	aa		ESTs, Weakly similar to RNG1_HUMAN RING1 PROTEIN [H.sapiens]
452	23834	AA956659	cc, General		EST
	1.0.45-		ļ.		ESTs, Moderately similar to C8
453	16425	AA956688	f,x		[M.musculus]
454	23847	AA956723	s		ESTs, Highly similar to Mi-2 protein
455	23852	AA956746	j,l,m,z		[H.sapiens]
456	5989	AA956907	g,s		ESTs, Highly similar to p162 protein [M.musculus]
456	5990	AA956907	General		ESTs, Highly similar to p162 protein [M.musculus]

TABLE 1:	SUMMARY				Atty , Docket No. 44921-5089W0 Doc. No. 1798697.1
Segvence ID No.	ldentifier	GenBenik Aee/ Ref. Seg ID	Model Gode	Gene Namo	Unigene Cluster Title
457	23957	AA957123	u,General		ESTs, Weakly similar to AF187065 1 p75NTR-associated cell death executor [R.norvegicus]
458	22357	AA957264	General		ESTs, Highly similar to hypothetical protein [H.sapiens]
450	23314	AA057270	g,l,m,p,v, cc, General		ESTs
459 460	23995	AA957270 AA957292	a,b		ESTs
				Libbarand tONA continue	ESTs, Moderately similar to SYS_HUMAN SERYL-TRNA
461	2702	AA957307	General	HHs:seryl-tRNA synthetase	SYNTHETASE [H.sapiens] ESTs, Highly similar to HIGH AFFINITY IMMUNOGLOBULIN EPSILON RECEPTOR GAMMA- SUBUNIT PRECURSOR
462	24040	AA957422	c		[R.norvegicus] ESTs, Highly similar to P3 MOUSE P3
463	12478	AA957554	m		PROTEIN [M.musculus]
464	21306	AA957811	٧		ESTs
465	24183	AA957889	t		ESTs
466	24178	AA957905	d		ESTs Highly similar to epsilon-COP
467	17034	AA963071	е		[M.musculus] ESTs, Weakly similar to AF187065 1 p75NTR-associated cell death
468	24053	AA963092	General		executor [R.norvegicus]
469	2767	AA963201	0		ESTs ESTs
470 471	2022 2126	AA963259 AA963488	g d		ESTs
472	24246	AA963703	b		ESTs, Highly similar to cell cycle protein p38-2G4 homolog [H.sapiens]
473	2195	AA963746	General		ESTs
474	19370	AA963797	i		ESTs
475	2282	AA964147	e		ESTs
476	2284	AA964152	×		EST
					ESTs, Highly similar to TGT_HUMAN QUEUINE TRNA-
478	2350	AA964368	g,General		RIBOSYLTRANSFERASE [H.sapiens] ESTs, Highly similar to ATRTC actin
479	18830	AA964496	aa		beta - rat [R.norvegicus]
480	2392	AA964541	b		EST
481	2395	AA964554	General		ESTs, Highly similar to U3 snoRNP associated 55 kDa protein [H.sapiens]
482	2410	AA964589	i,aa		EST
483 484	19145 2424	AA964613 AA964617	t g		ESTs ESTs
485	3107	AA964687	General		ESTs
486	2457	AA964752	q,t		EST
487	6778	AA964763	b		ESTs, Highly similar to DRIM protein [H.sapiens]
					ESTs, Weakly similar to T23337 hypothetical protein K05C4.2 -
489	2468	AA964807	<u>I</u>	Glutamate-cysteine ligase	Caenorhabditis elegans [C.elegans] Glutamate-cysteine ligase (gamma-
490	2469	AA964814	w	(gamma-glutamylcysteine synthetase), regulatory	glutamylcysteine synthetase), regulatory
491	12561	AA964815	General		ESTs
492	2326	AA964892	aa		ESTs, Highly similar to PROCOLLAGEN ALPHA 1(IV) CHAIN PRECURSOR [M.musculus]
493	21339	AA964962	General		ESTs, Highly similar to ABC1 MOUSE ATP-BINDING CASSETTE, SUB- FAMILY A, MEMBER 1 [M.musculus]
494	21339	AA964988	General		ESTs
495	12569	AA965023	g		ESTs

176	SUMMARY		11 (10.7) 12		Atty. Docket No. 44921-5089W6 Doc. No. 1798897.
Sequence ID Now!	Tellmed	Genbenk (Acc) Ref. Seq ID	Model Gode	Gene Name	Unigene Gluster Title
496	2583	AA965166	bb		ESTs, Moderately similar to inorganic pyrophosphatase [H.sapiens]
497	15885	AA965207	r		ESTs, Highly similar to KIAA0958 protein [H.sapiens]
499	2905	AA996727	b,l,m,u, General		ESTs
500 501	2915 2920	AA996782 AA996813	u,bb d		ESTs, Moderately similar to S27267 lamin A - rat [R.norvegicus]
501	2920	AA990013	aa,Gener		E318
502	19525	AA996856	al		EST
503	2984	AA997015	С		ESTs
504	2986	AA997028	General		ESTs
505	3145	AA997237	General		ESTs
506	19249	AA997342	m		ESTs
507	16883	AA997345	General		ESTs, Weakly similar to nitrilase homolog 1 [M.musculus]
508	12598	AA997362	s		ESTs, Moderately similar to LONN_HUMAN MITOCHONDRIAL LON PROTEASE HOMOLOG PRECURSOR [H.sapiens]
500	0.470				ESTs, Weakly similar to LIS1 MOUSE PLATELET-ACTIVATING FACTOR ACETYLHYDROLASE IB ALPHA
509 510	3470 3180	AA997374 AA997425	P		SUBUNIT [R.norvegicus]
511	3245	AA997608	General		ESTs, Weakly similar to PAI2 RAT PLASMINOGEN ACTIVATOR INHIBITOR-2, TYPE A [R.norvegicus]
512	3020	AA997656	t		ESTs, Moderately similar to T09071 SH3 domains-containing protein POSH - mouse [M.musculus] ESTs, Moderately similar to T30249
513	3269	AA997800	x,aa		cell proliferation antigen Ki-67 - mouse [M.musculus]
514	3288	AA997877	f	1	ESTs
515	23992	AA998164	k,x	Cyclin B1	Cyclin B1 ESTs, Moderately similar to
516	17470	AA998264	b		FLRE_HUMAN FLAVIN REDUCTASE [H.sapiens]
					ESTs, Weakly similar to BCL3_HUMAN B-CELL LYMPHOMA
517	3773	AA998356	General		3-ENCODED PROTEIN [H.sapiens]
518	19623	AA998422	General		ESTs, Highly similar to CGA2 MOUSE
519	3572	AA998516	×		CYCLIN A2 [M.musculus] ESTs, Moderately similar to CYCLIN- DEPENDENT KINASE INHIBITOR 1C
520	2782	AA998565	c i,r,w,		[M.musculus]
521	26119	AA998576	General		
522	22737	AA998660	aa		ESTs
523	3696	AA999030	е		ESTs, Moderately similar to AF132966 1 CGI-32 protein [H.sapiens]
524	3079	AA999169	k,x, General		ESTs
525	3081	AA999171	e,p,r	Signal transducer and activator of transcription 1	Signal transducer and activator of transcription 1
				HHs:guanine monphosphate	ESTs, Highly similar to GUAA_HUMAN GMP SYNTHASE
526	3082	AA999172	General	synthetase	[H.sapiens]
527	17337	AB000717 AB000778	k a	Phoshpolipase D gene 1	ESTs Phoshpolipase D gene 1
520					
528 529	1535	AB002406	k	RuvB-like protein 1	RuvB-like protein 1

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TABLE 1	SUMMARY				AMY. Docket No. 44921-5039WO
	6				Doc. No. 1793397.1
Seelieve Seelieve	9 	Conbank Acc Reil Seg ID	Model Gode	Gene Name	Uniçana Ciuster Tillo
10000	lucentimen #			Conte todates	Rattus norvegicus mRNA for
				9 T	carboxylesterase precursor, complete
531	4312	AB010635	c,i,j,k,y,z	HMm:DNA methyltransferase	cds ESTs, Highly similar to JE0378 DNA
532	21666	AB012214	k	(cytosine-5) 1	[R.norvegicus] Rattus norvegicus mRNA for G protein
533	15772	AB015645	g		coupled receptor, complete cds
					Rattus norvegicus MAP-kinase phosphatase (cpg21) mRNA,
534	1183	AF013144	h		complete cds
505	4500	AF045044	_		Rattus norvegicus NAC-1 protein
535	1582	AF015911	h,z		(NAC-1) mRNA, complete cds ESTs, Moderately similar to MY16
536	11483	AF020618	u,cc, General		MOUSE MYELOID DIFFERENTIATION PRIMARY RESPONSE PROTEIN MYD116 [M.musculus],Rattus norvegicus progression elevated gene 3 protein mRNA, complete cds
					Rattus norvegicus MHC class lb M4 (RT1.M4) pseudogene, complete
537	20295	AF024712	aa		sequence
					Rattus norvegicus chemokine CX3C
538 539	19077 23044	AF030358 AF034218	y,z General	hyaluronidase 2	mRNA, complete cds hyaluronidase 2
540	25178	AF035955	d	nyalaromado 2	Try and the decision of the second of the se
					Rattus norvegicus kidney injury
541	1564	AF035963	x,bb, General		molecule-1 (KIM-1) mRNA, complete
341	1304	AI 003300	Jonorai		Rattus norvegicus NonO/p54nrb
542	8426	AF036335	f		homolog mRNA, partial cds Rattus norvegicus homocysteine
					respondent protein HCYP2 mRNA,
543	21817	AF036537	k		complete cds
544	21145	AF038571	General	Solute carrier family 1 A1 (brain glutamate transporter)	Solute carrier family 1 A1 (brain glutamate transporter)
344	21143	A 03037 1	Ocheral	putative peroxisomal 2,4-	putative peroxisomal 2,4-dienoyl-CoA
545	22602	AF044574	General	dienoyl-CoA reductase	reductase
546	13464	AF047707	h	UDP-glucose:ceramide glycosyltransferase	UDP-glucose:ceramide glycosyltransferase
547	24024	AF052695	×	cell cycle protein p55CDC	cell cycle protein p55CDC
			i.		Rattus norvegicus trp1 beta variant
548	12259	AF061266	h	transient receptor protein 1	mRNA, complete cds Rattus norvegicus kidney-specific
549	4589	AF062389	y,z		protein (KS) mRNA, complete cds
				nuele seems seesmbly	Rattus norvegicus nucleosome assembly protein mRNA, complete
550	16007	AF062594	lt	nucleosome assembly protein 1-like 1	cds
	1.5557	1	1		Rattus norvegicus pyruvate
554	45704	A F060744	I		dehydrogenase phosphatase
551	15761	AF062741	u	 	isoenzyme 2 mRNA, complete cds Rattus norvegicus bithoraxoid-like
552	17426	AF073839	р		protein mRNA, complete cds
553	18615	AF074608	s	RT1 class lb gene	RT1 class lb gene Rattus norvegicus serine/threonine
554	15797	AF084205	f		protein kinase TAO1 mRNA, complete cds
			1		Rattus norvegicus 190 kDa ankyrin
555	12932	AF102552	s	ankyrin 3 (G)	isoform mRNA, complete cds ESTs, Highly similar to A49013 tumor
					cell suppression protein HTS1
556	18603	Al007649	x		[H.sapiens]
557 558	22733 22746	AI007668 AI007672	r		ESTs ESTs
559	24109	AI007072	General		ESTs

TABLE 18 S					Aiy. Docket No. 44921-5039W0 Doc. No. 1793397.
Sequence ID No.			Model Code	Cene Namo	Universe Cluster Title
					ESTs,ESTs, Highly similar to HS9B RAT HEAT SHOCK PROTEIN HSP
560	15848	A1007820	n,v	Hrs	90-BETA [R.norvegicus] Hrs
561 562	10108 6804	AI007857 AI007877	General	riis	ESTs
563	20099	A1007877	f,u		ESTs
564	11368	AI007948	d		ESTs, Weakly similar to T18778 hypothetical protein B0513.2b - Caenorhabditis elegans [C.elegans]
565	15849	A1008074	h		ESTs,ESTs, Highly similar to HS9B RAT HEAT SHOCK PROTEIN HSP 90-BETA [R.norvegicus]
566	3121	Al008160	General		ESTs, Moderately similar to AF15184 1 CGI-83 protein [H.sapiens] ESTs, Highly similar to Chain G, G
567	16646	AI008190	l t		Protein Heterotrimer Gi alpha 1 Beta 1 Gamma 2 With Gdp Bound
568	12683	A1008190	×		ESTs, Weakly similar to G2/MITOTIC SPECIFIC CYCLIN B1 [R.norvegicus]
					ESTs, Moderately similar to PIM1 RA PROTO-ONCOGENE SERINE/THREONINE-PROTEIN
569	22018	A1008309	b		KINASE PIM-1 [R.norvegicus] ESTs, Highly similar to
570	23917	AI008441	n		6PGD_HUMAN 6- PHOSPHOGLUCONATE DEHYDROGENASE, DECARBOXYLATIN [H.sapiens]
571	22599	AI008458	General		ESTs
572	22698	AI008578	p,General		ESTs
573	14405	AI008579	r,x		ESTs
574	4086	AI008629	x i,v,		ESTs, Moderately similar to JH0446 75K autoantigen [H.sapiens] ESTs, Weakly similar to heat shock
575	3808	AI008643	General		protein hsp40-3 [M.musculus] ESTs. Weakly similar to T29897
576	3931	Al008697			hypothetical protein F38A5.1 - Caenorhabditis elegans [C.elegans]
577	7785	AI008758	aa	Dipeptidyl peptidase 4	Dipeptidyl peptidase 4
					ESTs, Weakly similar to LONN_HUMAN MITOCHONDRIAL LON PROTEASE HOMOLOG
578	16701	Al008838	9		PRECURSOR [H.sapiens] ESTs, Weakly similar to CYSR RAT CYSTEINE-RICH PROTEIN 1
579	21789	A1008930	k	l	[R.norvegicus]
580	21895	AI008971	General		ESTs R.norvegicus mRNA encoding 45kDa
581	410	Al008974	i,aa, General		protein which binds to heymann nephritis antigen gp330 ESTs, Highly similar to BAG-family
582	21632	Al009167	General		molecular chaperone regulator-2 [H.sapiens]
583	21596	AI009168	General		ESTs
584	22801	Al009197	General cc,		ESTs ESTs, Highly similar to similar to human DNA-binding protein 5
585	11876	AI009321	General		[H.sapiens]
586	2506	AI009341	General		ESTs
587	6382	AI009362	General		ESTs
588	14370	AI009427	k		ESTs, Highly similar to Lmp10 proteasome subunit [M.musculus]

TABLE 1:	SUMMARY				A lly, Poeks No. 44921-5039WG Dog. No. 1793397.
Sequence ID No.	refilirebl	Cenbank Acc/ Ref. Seq ID	Model Code	Gene Name	Unigene Chester Title
				7 III. 33 January 1941 - 3 January 1941	ESTs, Highly similar to filamin
589 590	19275 4154	AI009460 AI009467	X		[H.sapiens]
590	3464	AI009407	g		ESTs
					ESTs, Highly similar to molybdopterin
592	3926	A1009592	e		synthase large subunit [M.musculus]
593	19358	AI009675	С		EST ESTs
594	22545	AI009747	g cc,		
595	15089	AI009752	General		ESTs
596	5458	AI009756	h	ALG-2 interacting protein 1	ALG-2 interacting protein 1
597	6844	AI009770	e,r,cc		ESTS
598	15627	AI009810	aa		ESTs, Highly similar to RS16_HUMAN 40S RIBOSOMAL PROTEIN S1 [R.norvegicus]
599	22619	Al009825	d		ESTs
600	7857	A1009898	j,l,m,z		ESTs ESTs
601 602	13259 21105	AI009946 AI010067	General		ESTS
002	21105	121010001	Seneral	Testis enhanced gene	2019
603	24627	AI010102	aa	transcript	Testis enhanced gene transcript ESTs, Moderately similar to
			_		YA00_HUMAN HYPOTHETICAL PROTEIN CGI-100 PRECURSOR
604	12716	Al010178	General		[H.sapiens]
605	18757	AI010216	aa	· · · · · · · · · · · · · · · · · · ·	ESTs ESTs, Weakly similar to claudin-7
606	2912	AI010220	aa, General		[R.norvegicus]
607	3316	Al010220	t		ESTs
608	15644	AI010257	General		R.norvegicus mRNA for histone H3.3
609	657	AI010262	b		Rattus norvegicus mRNA for inetrleukin-4 receptor (membrane-bound form), complete cds
610	3271	Al010303	b		ESTs
					ESTs, Moderately similar to erythroblast macrophage protein EMP
611	11081	Al010407	c,s,t,		[H.sapiens]
612	16521	AI010470	General	Ceruloplasmin (ferroxidase)	Ceruloplasmin (ferroxidase)
613	6927	Al010542	General		ESTs
614	17524	AI010568	a,j,y, General	Growth hormone receptor	Growth hormone receptor
615	6946	AI010642	n		ESTs
616	23509	AI010962	aa		ESTs, Highly similar to SDP3 [M.musculus]
617	6044	AI010302	t		ESTs
618	13855	Al011265	0		ESTs
619	21779	AI011380	cc		ESTs
621	12534	AI011460	СС		ESTs
622	12629	AI011492	e,f		ESTs, Moderately similar to HYA22 [H.sapiens]
	12020				ESTs, Weakly similar to B Chain B, Solution Structure Of The C-Terminal Negative Regulatory Domain Of P53 In A Complex With Ca2+-Bound
623	735	AI011560	f		S100b(Bb) [R.norvegicus] ESTs, Moderately similar to LMA5
624	3941	AI011598	General		MOUSE LAMININ ALPHA-5 CHAIN [M.musculus] ESTs, Weakly similar to JE0360
625	17550	AI011607	j,General		gamma-Butyrobetaine hydroxylase [H.sapiens] ESTs, Weakly similar to I(3)S12
	1	AI011634	1.		protein [D.melanogaster]
626	10636		le		

TABLE 18 S	WIXIMARY)				AMy, Docket No. 44921-5089W0
					Dos. No. 1793397.1
Sequence ID No.	0-0	Conbank Acel Ref. Seq ID	Model	Gane Name	Unigene Gluster Tille
I B. IVO.	ncenuner	Kensedin	COUC		ESTs, Weakly similar to SFR5 RAT
					SPLICING FACTOR,
					ARGININE/SERINE-RICH 5
628	16112	AI011706	h		[R.norvegicus]
600	12254	A1044757			ESTs, Weakly similar to A35902 Fc
629 630	13354	AI011757 AI011799	cc		gamma [R.norvegicus] IESTs
	1	7.101.11.00			ESTs, Highly similar to AF151842 1
631	18684	AI011812	t		CGI-84 protein [H.sapiens]
632	4205	AI011982	b		ESTs
633	6518	AI012114	General	•	ESTs, Moderately similar to R29425 1 [H.sapiens]
634	17407	AI012114 AI012145	General		ESTs
	112	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,			ESTs, Weakly similar to PPP5 RAT
635	13093	AI012177	r		SERINE/THREONINE PROTEIN PHOSPHATASE 5 [R.norvegicus]
					ESTs, Moderately similar to
636	15395	Al012216	f		Y33K_HUMAN HYPOTHETICAL 33.4 KDA PROTEI [H.sapiens]
	1				
					ESTs, Weakly similar to S70484 RS43
637	21796 3981	AI012221	d,General i,General		protein - rat (fragment) [R.norvegicus] IESTs
638 639	6606	Al012235 Al012308	i,r		ESTs
000	0000	A1012300	','		ESTs, Highly similar to NHPX RAT
					NHP2/RS6 FAMILY PROTEIN
640	3417	AI012337	w		YEL026W HOMOLOG [R.norvegicus]
	04000	A10400E0	b,t,		EST-
641 642	24200 7471	AI012356 AI012379	General cc		ESTs ESTs
643	7247	AI012438	g		ESTs
644	7127	AI012464	p,General		ESTs
					ESTs, Weakly similar to T26998 hypothetical protein Y48B6A.6 -
645	3304	AI012471	ь		Caenorhabditis elegans [C.elegans]
646	2311	AI012485	aa		ESTs
				glutathione S-transferase, pi	
647	20817	AI012589	g,n,q	2	glutathione S-transferase, pi 2
648 649	3493 8975	AI012590 AI012613	v,General General		ESTs ESTs
043	03/3	A1012013	Certeral		ESTs, Highly similar to unknown
650	11335	AI012619	j		[H.sapiens]
651	21409	AI012637	General		ESTs
					ESTs, Moderately similar to AF151834
652	8015	AI012638	aa		1 CGI-76 protein [H.sapiens]
002	10010	7410 12000			ESTs, Highly similar to RS20_HUMAN
					40S RIBOSOMAL PROTEIN S2
653	8476	AI012647	w		[R.norvegicus]
654	4232	A1012059	e,p, General		ESTs
655	23128	AI012958 AI013011	General		ESTs
656	20086	AI013260	General	lamin	lamin
· · · · · · · · · · · · · · · · · · ·					ESTs, Highly similar to GLIA DERIVED NEXIN PRECURSOR
657	11969	AI013273	k		[R.norvegicus]
658	26147	AI013387	aa		
659	8815	AI013437	p		ESTs
660	19722	AI013508	k		Rattus norvegicus Hsp70 binding protein HspBP mRNA, complete cds
661	6674	AI013568	General		ESTs
662	23145	AI013647	o,t		ESTs
663	15130	AI013676	w		ESTs

				-71-	
TABLE 1: S					Aiiy. Docket No. 44921-503900 Doc. No. 1793897
Sequence	lidentifier:	Genbenk Acci. Ref. Sog ID	Model Gode	Gene Name	Unicene Cluster Tille
Section 1	3) has been dead and a supplied that a supplied to the supplined to the supplied to the supplied to the supplied to the suppli	A SAME AND		and the second s	ESTs, Moderately similar to BMP6 RAT BONE MORPHOGENETIC PROTEIN 6 PRECURSOR
664	7274	Al013715	aa		[R.norvegicus] ESTs, Highly similar to KIAA1102
665	7276	AI013730	е		protein [H.sapiens]
666	7278	Al013738	y,z,aa		ESTs
667	22592	AI013740	s,x,bb, General		ESTs, Highly similar to proteolipid protein 2 [M.musculus]
668	16584	Al013765	w	Arrestin, beta 2	Arrestin, beta 2
					ESTs, Highly similar to T27225 ADP- ribosylation factor Y57G11C.13 [similarity] - Caenorhabditis elegans
669	24143	AI013804	j,I		[C.elegans]
670	15928	AI013829	a,General		ESTs
671	21950	AI013861	l _i	3-hydroxyisobutyrate dehydrogenase	3-hydroxyisobutyrate dehydrogenase
672	3260	AI013875	t	deriydrogenase	ESTs
					ESTs, Moderately similar to MSSP
673	2708	AI013882	d,q		[M.musculus]
674	8585	AI013886	p,r,t,	<u> </u>	ESTs ESTs, Weakly similar to CIRP
675	7299	Al013911	General		[R.norvegicus] Rat ankyrin binding glycoprotein-1
676	15904	AI013971	General		related mRNA sequence
677	12781	AI014023	w		ESTs, Moderately similar to R32184 [H.sapiens]
					Rattus norvegicus mRNA for beta- carotene 15,15'-dioxygenase,
678	19372	AI014135	aa		complete cds ESTs, Highly similar to hypothetical
679	4241	AI014140	w		protein [H.sapiens]
680	15247	Al014169	c,u		Rattus norvegicus clone N27 mRNA ESTs, Moderately similar to mitogen-
C04	7245	A1020024	<u></u>		activated protein kinase kinase kinas
681 682	7315 16631	AI028831 AI028856	n General		6 [H.sapiens]
683	23297	AI028953	x		ESTs, Highly similar to S55054 Sm protein G [H.sapiens]
684	11326	Al028935	b		ESTs
685	2866	AI029058	n,y		ESTs
686	12812	Al029126	General		ESTs
687 688	17602 7392	AI029156 AI029185	p aa		ESTs EST
689	6517	AI029185	d,k,x		ESTs
690	7639	AI029292	b		ESTs
					ESTs, Highly similar to CB80_HUMA 80 KDA NUCLEAR CAP BINDING
691	3874	AI029428	i,General		PROTEIN [H.sapiens]
692 693	12819 7452	AI029437	f r		ESTs ESTs
694	7493	AI029466 AI029608	b		ESTS
696	7537	AI029829	o,General		ESTs
697	2310	AI029969	v		ESTs
698	7585	A1030023	X b n		ESTs ESTs
699 700	7586 14492	AI030024 AI030091	b,n cc		ESTs
701	10673	AI030134	f		ESTs, Weakly similar to ankyrin [R.norvegicus]
702	7615	AI030163	o,r		ESTs
703	2370	AI030179	General		ESTs
704	7681	A1030449	n		ESTs, Moderately similar to methyltransferase related protein [M.musculus]

TABÚE 18 {			11.00		Atty. Docket No. 44921-503900 Doc. No. 1793397.
Sequence ID (No.	lieniiler	GenBank Acel Ref. Seq (D	Model Gode	Cene Name	Uniter: Cheter Tile
705	11559	AI030472	General	and the second s	ESTs
	11000	711000112			Rattus norvegicus nucleosome
					assembly protein mRNA, complete
706	7665	AI030668	t,bb		cds
707	24222	AI030704	k		ESTs
708	10740	Al030743	h		EST
709	10742	AI030773	е		EST
744	1,0,100	41000000	Camana1		ESTs, Moderately similar to
711 712	16169	AI030932 AI030991	General		adipophilin [H.sapiens]
/12	19527	Alususai	 		ESTs, Highly similar to SX17 MOUSE
					TRANSCRIPTION FACTOR SOX-17
713	22614	AI031004	l _r		[M.musculus]
7 13	22014	7.1001004		<u> </u>	ESTs, Highly similar to CLPP MOUS
					PUTATIVE ATP-DEPENDENT CLP
					PROTEASE PROTEOLYTIC
					SUBUNIT, MITOCHONDRIAL
714	3167	AI031012	е		PRECURSOR [M.musculus]
715	5350	Al043611	а		ESTs
716	7858	AI043654	t		EST
717	10784	AI043678	d		EST
					ESTs, Weakly similar to T27134
					hypothetical protein Y53C12B.2 -
718	9180	AI043694	aa		Caenorhabditis elegans [C.elegans]
	İ			HHs:phosphoribosyl	
		1		pyrophosphate	Rattus norvegicus mRNA for
719	7867	AI043695	aa	amidotransferase	amidophosphoribosyltransferase
720	7584	AI043724	General		ESTs
		1.0.0700			ESTs, Highly similar to AF151810 1
721	7895	AI043768	e Caracal		CGI-52 protein [H.sapiens]
722	7903	AI043805	General	· · · · · · · · · · · · · · · · · · ·	ESTs, Weakly similar to ELL MOUSE
					RNA POLYMERASE II ELONGATION
723	7913	AI043849	СС		FACTOR ELL [M.musculus]
724	3899	AI043904	100	 	ESTs
725	6766	AI043914	f		ESTs
	10.00		g,I,m,		
726	10818	AI043990	General		ESTs
727	7956	AI044018	f		EST
728	5393	AI044170	р		EST
729	5398	AI044177	q		EST
					ESTs, Weakly similar to AF121893 1
					sequence-specific single-stranded-
730	5425	AI044237	a,d		DNA-binding protein [R.norvegicus]
					ESTs, Weakly similar to putative
					peroxisomal 2,4-dienoyl-CoA
731	8692	AI044247	r	<u> </u>	reductase [R.norvegicus]
732	5430	AI044253	1 -	 	EST
722	E461	A1044220	g,p, General		 ESTs
733	5461	A1044338	General		ESTS
734 735	5464 3359	Al044345 Al044347	aa		ESTS
100	3333	ורטדרטורן	aa		Rat (clones rLG[08,14,25]) interleukir
737	2695	AI044396	Ь		6 signal transducer mRNA sequence
73 <u>7</u> 738	5494	AI044396 AI044425	General		ESTs
740	9882	AI044588	j,m	 	ESTs
741	5575	AI044688	g	 	ESTs
742	2348	AI044794	General		ESTs
	1		1 2		ESTs, Weakly similar to AF165892 1
		1			RNA-binding protein SiahBP
743	18205	AI044836	n		[R.norvegicus]
744	5626	AI044864	u		ESTs
745	5630	AI044869	f		ESTs
					ESTs, Moderately similar to AF15187
746	5634	AI044883	General	1	1 CGI-115 protein [H.sapiens]

TABLE 18	SUMMARY				Atty. Docket No. 44921-5089V Doc. No. 1798697
© 170° gednevoe ¹	ldeniffer	Conbook Ace/ Rof. Seq ID	Model (606) (6	ene Name	Unigene Gueter Title
			B and who we have a second was a		ESTs, Moderately similar to
747	4047	A1044947	l,m		dJ1183I21.1 [H.sapiens]
748 749	5654 5684	AI044976 AI045056	r		IESTs
45	3004	A1043036			ESTs, Highly similar to BGAL MOUS BETA-GALACTOSIDASE
750	19235	AI045074	General i.aa.		PRECURSOR [M.musculus] ESTs, Moderately similar to HEM45
751	5689	AI045075	General		[H.sapiens] ESTs, Moderately similar to AF1188
752	5711	AI045151	General		1 citrin [H.sapiens] ESTs, Weakly similar to TVRTK6
	1		1_		ribosomal protein S6 kinase
753 754	19237 19964	AI045153 AI045161	C		[R.norvegicus]
754 755	5735	AI045161 AI045223	 		IESTs
			a,General		ESTs
756 757	5474 5811	Al045477 Al045502	d.e		ESTs
758	5819	AI045537	General		IESTs IESTs
759	5839	AI045594	i		ESTs
760	6808	AI045600	s		ESTs, Highly similar to S30034 translocating chain-associating membrane protein [H.sapiens]
761	17755	AI045608	у		ESTs
763	10020	AI045632	a		ESTs
764	5855	AI045669	General		ESTs
765	5881	AI045789	i	· <u>· · · · · · · · · · · · · · · · · · </u>	ESTs, Weakly similar to T12540 hypothetical protein DKFZp434J214 [H.sapiens] ESTs, Moderately similar to S64732
766	5897	AI045862	General		scaffold attachment factor B [H.sapiens]
767	5900	AI045866	y,z		ESTs
			o,t,		ESTs, Weakly similar to B48013 proline-rich proteoglycan 2 precurso
768	7540	AI045882	General		parotid - rat [R.norvegicus]
769	5329	Al045970	Р		ESTs
770	15093	AI058285	d		ESTs
771	8002	AI058304	+		EST EST
772	8017	AI058341	С		ESTs, Weakly similar to T46465 hypothetical protein
773	6828	AI058359	General		DKFZp434A0530.1 [H.sapiens]
774	8177	AI058603	aa		ESTs
775	3090	AI058730	aa		ESTs
776	10093	AI058746	g General		ESTs ESTs
777 778	8143 18659	AI058759 AI058762	General		ESTS ESTS
779	8163	AI058762 AI058837	laa		ESTs
780	4789	AI058889	General		ESTs
781	8221	AI059061	General		ESTs
782	10159	AI059147	d		EST
783	8245	AI059154	b		ESTs, Weakly similar to unnamed protein product [H.sapiens]
784	8283	AI059290	n		ESTs
785	8314	AI059386	g,General		ESTs
786	10200	A1059386 A1059444	ig,General		ESTs
700	10200	A1059444			ESTS, Weakly similar to EGF RAT PRO-EPIDERMAL GROWTH FACTOR PRECURSOR
787	8347	AI059519	s		[R.norvegicus] Rattus norvegicus transitional endoplasmic reticulum ATPase
788	18359	AI059675	n		mRNA, complete cds

TABUE 1:	SUMMARY				Atty. Docket No. 44924-5039WG Doc. No. 1793397.
Sequence ID No.	lientifier	Conbank Ace Ref. Seq ID	Model : Gode	Cene Name	Unigeno Gluster Title
789	10281	AI059947	b,t		EST
790	8494	AI059968	aa		ESTs
					ESTs, Weakly similar to TNRC
					MOUSE LYMPHOTOXIN-BETA
				•	RECEPTOR PRECURSOR
791	8495	AI059971	General		[M.musculus] ESTs, Moderately similar to KIAA097
700	8496	A1050074	General		protein [H.sapiens]
792	0490	AI059974	General		ESTs, Weakly similar to CGI-142
793	10289	AI060053	li		hypothetical protein [H.sapiens]
794	8548	AI060176	k	-	ESTs
795	8565	AI060236	t		EST
796	18322	AI060279	i,y,z		ESTs
797	8745	AI069939	r		ESTs
					ESTs, Highly similar to rer
798	8785	AI070067	lo		[M.musculus]
					ESTs, Weakly similar to 2104282A
799	17506	AI070068	cc		Gadd45 gene [R.norvegicus]
					ESTs, Weakly similar to NUCL RAT
800	9067	A1070087	General		NUCLEOLIN [R.norvegicus]
					ESTs, Moderately similar to CGI-97
801	3551	Al070122	e		protein [H.sapiens]
					ESTs, Moderately similar to GLMB
					RAT GLIA MATURATION FACTOR
802	4967	AI070179	k		BETA [R.norvegicus]
803	18	AI070195	General	:	ESTs, Moderately similar to AF13295 1 CGI-20 protein [H.sapiens]
					ESTs, Moderately similar to ARVC_HUMAN ARMADILLO REPEAT PROTEIN DELETED IN VELO-CARDIO-FACIAL SYNDROME
804	24197	AI070314	General		[H.sapiens]
805	8869	AI070330	r		ESTs
806	8874	A1070336	b,cc		ESTs
807	10417	AI070410	m		ESTs ESTs, Moderately similar to T08664 Toll protein-like receptor
808	8901	AI070419	aa		DKFZp547l0610.1 [H.sapiens]
			l,p,	177	
809	14424	AI070421	General		ESTs
810	10434	AI070497	General		ESTs
811	8927	AI070523	V		ESTs
812	8946	AI070611	q		ESTs
813	8950	AI070621	w		ESTs
814	8972	AI070673	General		ESTs
815	8981	AI070715	bb		EST
816	26184	A1070784	i,l		ESTs, Weakly similar to hypothetical
817	3007	Al070824	w	<u> </u>	protein [H.sapiens]
818	8999	A1070839	р		ESTs
819	10477	AI070868	e,f	bone morphogenetic protein 1 (procollagen C-proetinase)	bone morphogenetic protein 1 (procollagen C-proetinase)
820	24301	Al070911	k	1	ESTs
OLU		Al071024	General		EST
	8721		×		ESTs
821	9212	AI071098	1^		
821 822		AI071098 AI071137	c		Rat mRNA for cdc25B, complete cds
821 822 823	9212				EST
821 822 823 824	9212 1831 11005	Al071137 Al071139	r		EST ESTs, Highly similar to HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEIN G
821 822 823	9212 1831	AI071137			EST ESTs, Highly similar to HETEROGENEOUS NUCLEAR

5 (A) (A)	SUMMARY				Ally, Docket No. 44921-5039W0 Doc. No. 1793397.1
Sequence ID (Vo.	e lidentifier	GenBank Acc/ Ref. Seq ID	Model Code	Cene Name	Unigene Cluster Title
				HHs:serine palmitoyltransferase, long	ESTs, Highly similar to JC5180 serine
828 829	16058 11057	AI071490 AI071509	General f,o	chain base subunit 2	C-palmitoyltransferase [M.musculus] ESTs
					ESTs, Weakly similar to SYBSR threonine synthase (EC 4.2.99.2) - yeast (Saccharomyces cerevisiae)
831	5695	AI071566	bb		[S.cerevisiae]
832	9671	AI071568 AI071578	W		ESTs, Moderately similar to NEURONAL PROTEIN 3.1 IM.musculus
834	9673	AI071581	General		ESTs
835	9699	Al071646	General		ESTs
837	9799	AI072008	q,y,z		ESTs
838	9808	AI072050	d		ESTs
839	22796	Al072213	General		ESTs
840	9271	AI072405	V		ESTs
841	10869	AI072425	w		ESTs ESTs, Weakly similar to S70484 RS43
842	21797	AI072439	General		protein - rat (fragment) [R.norvegicus]
843	9306	AI072521	r		ESTs
844 845	9312 10893	AI072550 AI072559	X		ESTs EST
846	1501	Al072539 Al072634	cc,Gener		Rattus norvegicus cytokeratin-18 mRNA, partial cds
847	6548	AI072658	General		ESTs
					ESTs, Highly similar to JE0170 dnaJ heat shock protein MCG18 - mouse
848	9363	A1072695	d		[M.musculus] ESTs, Moderately similar to LMG2 MOUSE LAMININ GAMMA-2 CHAIN
850	9409	AI072841	n		PRECURSOR [M.musculus]
851	9410	AI072842	w		ESTs
852	9468	AI073021	General		ESTs
853	9518	AI073223	f		EST ESTs, Weakly similar to CAH2 RAT CARBONIC ANHYDRASE II
854	11183	AI100768	t	HHs:carbonic anhydrase VIII	
855	9190	AI100835	е		ESTs
856	2029	AI100842	р		ESTs
857	5687	AI101006	e		ESTs Rat metallothionein-2 and
858	15192	Al101099	g,cc		metallothionein-1 genes, complete cds ESTs, Highly similar to ATPK MOUSE ATP SYNTHASE F CHAIN,
859	17399	AI101157	0		MITOCHONDRIAL [M.musculus] ESTs, Weakly similar to S46930
860	9339	Al101160	l,m,o		teg292 protein - mouse [M.musculus] ESTs, Weakly similar to AIF-C1
861	6321	AI101256	General		[R.norvegicus] ESTs, Highly similar to GDIS MOUSE RHO GDP-DISSOCIATION
862	5421	AI101270	С		INHIBITOR 2 [M.musculus] ESTs, Highly similar to ERM_HUMAN
					ETS-RELATED PROTEIN ERM
863	11910	AI101323	General		[H.sapiens]
864	23140	AI101608	Coporal		ESTs
865	4119	AI101901	General		ESTs ESTs, Weakly similar to TRBP
866	16324	AI102009	b		MOUSE PROTAMINE-1 RNA BINDING PROTEIN [M.musculus]
867	18642	AI102023	o		ESTs, Moderately similar to unknown [H.sapiens]

TABLE 1:	SUMMARY				/ Aliy. Docket No. 44921-5039WC Doc. No. 1793397.
eguence Sequence	Identifier	GenBenk Ace/ Ref. Seq ID	Model Code	Gene Name	Unitgene Gluster Mile
868	19373	Al102044	а	Drosophila polarity gene (frizzled) homologue	Rattus norvegicus mRNA for beta- carotene 15,15'-dioxygenase, complete cds
869	7051	AI102055	h		Rattus norvegicus clone ZG52 mRNA sequence
870	6544	AI102064	С		ESTs, Weakly similar to AF147718 1 glycine decarboxylase [R.norvegicus]
871	10227	AI102248	w		ESTs ESTs
872	23849	AI102318	e,q		Rattus norvegicus liver cytochrome c
873	11954	AI102505	g,j,s	HMm:cytochrome c oxidase, subunit VIIIa	oxidase subunit VIII (COX-VIII) mRNA, 3' end of cds
874	2125	AI102519	c,k		ESTs, Moderately similar to DAP12 [M.musculus]
875	5967	AI102520	v		ESTs, Moderately similar to AF16158 1 GABA-A receptor-associated protein [R.norvegicus]
		-			ESTs, Moderately similar to AF16158 1 GABA-A receptor-associated protein
875	5969	AI102520	p,w		[R.norvegicus]
876	11563	AI102560	General		ESTS
877	15190	AI102562	b,g,n,p,v		Rat metallothionein-i (mt-1) mrna EST, Weakly similar to A60716 somatotropin intron-related protein
878	19769	AI102570	bb		RDE.25 - rat [R.norvegicus] ESTs, Highly similar to I49523 Mouse
					primary response gene B94 mRNA,
879	22487	AI102578	General		3'end - mouse [M.musculus]
880 881	19011 23837	AI102618 AI102620	General		ESTs ESTs
001	23837	A1102620	q,t	solute carrier family 20	2018
882	23538	AI102727	g,General	(phosphate transporter), member 1	solute carrier family 20 (phosphate transporter), member 1
883	17234	AI102741	С	metalloproteinase 3	Tissue inhibitor of metalloproteinase 3
884	5891	AI102745	k	motalioprotolilace s	ESTs
885	6796	AI102753	General		ESTs
886	8837	AI102849	о,р		ESTs
887	15861	AI102868			ESTs, Weakly similar to phosphoserine aminotransferase [H.sapiens]
888	3533	Al102877	g		ESTs
889	13222	AI102977	General		ESTs, Highly similar to PCAF associated factor 65 beta [H.sapiens]
890	6806	AI103018	o,u		ESTs
891	10659	Al103059	w,cc, General		ESTs
					ESTs, Highly similar to ATPK MOUSE ATP SYNTHASE F CHAIN,
892	17400	Al103097	e		MITOCHONDRIAL [M.musculus]
893	3584	AI103106	x,aa		ESTs ESTs
894	13298	AI103143	1		ESTs, Weakly similar to UBC2_HUMAN UBIQUITIN- CONJUGATING ENZYME E2-17 KD
895	15981	AI103150	i,x		[R.norvegicus] ESTs, Highly similar to AF151893 1
896	3475	AI103245	w .		CGI-135 protein [H.sapiens]
898	23619	AI103314	р		ESTs ESTs, Moderately similar to T26785 hypothetical protein Y40B1B.7 -
899	24181	AI103320	le		Caenorhabditis elegans [C.elegans]
901	4355	AI103410	General		ESTs
902	7622	AI103472	General		ESTs
903	20918	AI103552	n		ESTs
904	21579	AI103572	General		ESTs

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TABLE 18	SUMMARY -				Atty. Docket No. 44921-5089W0
			PARTY NUMBERS OF PERSONS		Doc. No. 1798397.1
Sequence ID No.		Conbank Ace.	Model	Ceme Name	Unigene Clusier Tille
פאש	: Identifiter	Kete Sed In	Codex	Gene Name	
205	2222	A1402624	0		ESTs, Highly similar to RIE2 [M.musculus]
905	2222	AI103631	 		ESTs, Highly similar to sarcosine
906	2752	AI103641	e		dehydrogenase [R.norvegicus]
907	4856	AI103708	i		ESTs
908	8990	AI103719	l,m,y,z		
909	15942	AI103738	r		ESTs
040	22005	A1402020	o Conorol		ESTs
910 911	22885 15853	AI103828 AI103841	e,General	Complement component 4	Complement component 4
911	13633	A1103041	 	HHs:ubiquinol-cytochrome c	Complement component 4
				reductase. Rieske iron-sulfur	Rat Rieske iron-sulfur protein mRNA,
912	15050	AI103911	j,y	polypeptide 1	complete cds
913	12376	AI103939	u		ESTs
					ESTs, Weakly similar to AF151109 1
					putative BRCA1-interacting protein
914	22271	A1103947	о,у		[H.sapiens]
				HMm:RIKEN cDNA	ESTs, Highly similar to COXG MOUSE CYTOCHROME C OXIDASE
915	20833	AI104035	f,q	2010000G05 gene	POLYPEPTIDE VIB [M.musculus]
916	7010	Al104099	w	20,00000000 go.i.c	ESTs
917	22101	AI104251	General		ESTs
918	22833	Al104258	General		ESTs
					ESTs, Highly similar to translation
919	22211	Al104279	g,m		initiation factor eIF6 [M.musculus]
920	10720	Al104296	<u> </u>		ESTs
921	15416	Al104340	-		ESTs ESTs
922	10991	Al104342	а		ESTs, Highly similar to ATRTC actin
923	18831	AI104357	p		beta - rat [R.norvegicus]
924	7223	AI104337	e		ESTs
	1	1	1	Cytochrome c oxidase	Cytochrome c oxidase subunit VIa
925	23574	Al104520	e,g,s	subunit VIa (liver)	(liver)
					ESTs, Weakly similar to
	1	1			NADH:ubiquinone oxidoreductase B17
926	18509	Al104528	 9		subunit [H.sapiens] ESTs
927	11680	Al104605	 ' 		ESTs, Weakly similar to RENAL
					TRANSCRIPTION FACTOR KID-1
928	12342	AI104658	lw		[R.norvegicus]
					Rat mitochondrial succinyl-CoA
					synthetase alpha subunit (cytoplasmic
929	23689	Al104685	r		precursor) mRNA, complete cds
					ESTs, Moderately similar to T50611
930	15277	A1404924	0.00		hypothetical protein DKFZp434H2035.1 [H.sapiens]
930	15377	AI104821	o,cc		ESTs, Moderately similar to
					meningioma-expressed antigen 11
931	22957	AI104897	General		[H.sapiens]
				HHs:ATP synthase, H+	
				transporting, mitochondrial	Rattus norvegicus delta subunit of
932	18451	AI104953	o,s	F1 complex, delta subunit	F1F0 ATPase gene, complete cds
022	24275	A1404070	n Conoral		ESTs, Moderately similar to nucleolar
933	24375	Al104979	n,General		protein p40 [H.sapiens] ESTs, Moderately similar to
					SCOT_HUMAN SUCCINYL-COA:3-
					KETOACID-COENZYME A
					TRANSFERASE PRECURSOR
934	18278	AI105080	bb		[H.sapiens]
935	2196	AI105243	g		ESTs TO 1044
			1	Ī	ESTs, Weakly similar to T21641
			h.		
936	5199	AI105272	bb, General		hypothetical protein F32B6.2 - Caenorhabditis elegans [C.elegans]

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VABUE 1;					Ally, Docket No. 44921-5039 Doc. No. 179839		
Sequence ID No.	ldeniiiler	Coneank Accept Ref. Seque	Model Code	Gene Name	Unigane Gluster Title		
					ESTs, Weakly similar to T19707		
938	7700	AI105383	cc, General		hypothetical protein C34C6.5 - Caenorhabditis elegans [C.elegans		
939	13343	AI105398	u	·	ESTs		
940	22931	AI105417	e,General		ESTs, Moderately similar to NEURONAL PROTEIN 3.1 [M.musculus]		
	00500			HMm:glutaryl-Coenzyme A	ESTs, Highly similar to GCDH MOUSE GLUTARYL-COA DEHYDROGENASE PRECURSOR		
941	23596	AI105435	bb	dehydrogenase	[M.musculus] ESTs, Moderately similar to DHSD_HUMAN_SUCCINATE		
942	15893	AI105465	o		DEHYDROGENASE [H.sapiens]		
943	12660	Al111492	С		ESTs		
944	4479	AI111599	General		ESTs ESTs, Highly similar to H33_HUMA		
945	24211	AI111853	k		HISTONE H3.3 [R.norvegicus] ESTs, Weakly similar to FKB5		
946	2539	AI111960	r		MOUSE 51 KDA FK506-BINDING PROTEIN [M.musculus]		
947	5729	AI111990	k		EGF-CONTAINING FIBULIN-LIKE EXTRACELLULAR MATRIX PROTEIN 1 PRECURSOR (FIBULI 3) (FIBL-3) (T16 PROTEIN) [R.norvegicus]		
948	4049	AI112012	i,q,u, General		Rattus norvegicus osteoactivin mR complete cds		
949	12908	Al112043	i		ESTs		
950	20041	AI112161	t		ESTs		
951	12937	AI112462	General		ESTs		
952	3713	Al112571	b		ESTs		
953	12921	Al112636	General		ESTs, Moderately similar to UDP_HUMAN URIDINE PHOSPHORYLASE [H.sapiens]		
954	12965	Al112926	General		ESTs		
955	7499	Al112986	General		ESTs ESTs, Moderately similar to megakaryocyte stimulating factor		
956	4969	Al136305	f		[H.sapiens] ESTs, Highly similar to BC-2 protei [H.sapiens]		
957	11817	AI136295			ESTs, Weakly similar to JC4975 plexin 2 precursor - mouse		
959 960	11165	AI136372 AI136460	cc		[M.musculus] ESTs		
961	12782	Al136493	k		ESTs		
962	6850	Al136665	h	ecto-apyrase	ecto-apyrase		
963	20920	Al136891	p,v	butyrate response factor 1	butyrate response factor 1		
964	6552	Al137062	0		ESTs, Highly similar to 6.2 kd prote [H.sapiens]		
965	22722	Al137211	ļi .		ESTs Highly similar to synetaral		
966 967	13111 15969	Al137224 Al137302	o,General		ESTs, Highly similar to oxysterol- binding protein [M.musculus] ESTs		
968	14349	Al137302	d		ESTs		
969	9166	Al137406	General		ESTs ESTs, Weakly similar to ZF37_RAT		
070	0525	A1127516			ZINC FINGER PROTEIN 37 (ZFP-3		
970 971	9525 6638	AI137516 AI137579	General		ESTs		
972	7414	Al137586	General		ESTs, Highly similar to IMB3_HUM IMPORTIN BETA-3 SUBUNIT [H.sapiens]		
973	11321	Al137752	z		ESTs		
974	23473	AI137932	Ti Ti	i	ESTs		

santaun Sings.	SUMMERT				Aiiy, Docket No. 44921-5039446 Doc. No. 1793397.
Sequen c e) ID No.	ldeniiier	Cenbenk Acel * Ref. See ID	Model Gode	Gene Name	Unigene Civeter Title
975	13158	Al138024	Destruction of the last		ESTs
313	13130	71130024	- -	UDP-glucose:ceramide	UDP-glucose:ceramide
976	13467	AI138034	СС	glycosyltransferase	glycosyltransferase
977	11377	Al138105	v		ESTs
978	6790	Al144801	d,h		ESTs
979	6506	Al144919	j,l,y		ESTs
980	8027	A1144958	i		ESTs
982	14458	AI145095	General		ESTs
983	7476	Al145202	9		ESTs
			L		ESTs, ESTs, Weakly similar to GTP-
984	17545	AI145384	e		binding protein [H.sapiens] ESTs
985	17479	AI145385	r		ESTs
986	4194	AI145387	 '		ESTs, Weakly similar to T31511
			1		hypothetical protein Y116A8C.9 -
987	8634	Al145722	g		Caenorhabditis elegans [C.elegans]
901	10034	A1143722	9		ESTs, Weakly similar to T21659
			1		hypothetical protein F32D8.4 -
988	8339	Al145761	v.General		Caenorhabditis elegans [C.elegans]
	13000		,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,		ESTs, Highly similar to pseudouridine
989	2059	AI146005	h,General		synthase 1 [M.musculus]
	1	1	†		Rattus norvegicus small zinc finger-
990	23224	AI146033	0		like protein (TIM9a) mRNA, partial cds
				branched chain keto acid	
			1	dehydrogenase E1, beta	branched chain keto acid
991	5232	Al168942	bb	polypeptide	dehydrogenase E1, beta polypeptide
992	18472	Al168975	u		ESTs
992	18473	Al168975	u		ESTs
993	13235	Al169020	r		ESTs
			o,y,		
994	11618	Al169115	General		ESTs
					ESTs, Weakly similar to T23206
005	47000	1400444			hypothetical protein K01H12.1 -
995	17386	AI169144	0		Caenorhabditis elegans [C.elegans] ESTs, Weakly similar to HP33
996	10984	AI169156	o,u		[R.norvegicus]
997	8205	Al169176	e e		ESTs
331	0203	A1103170	-		ESTs, Highly similar to RADIATION-
					INDUCIBLE IMMEDIATE-EARLY
998	12979	AI169177	е		GENE IEX-1 [M.musculus]
-	1.2070	7.1.100 11.7			ESTs. Highly similar to A47318 RNA-
					binding protein Raly - mouse
999	2607	Al169211	c		[M.musculus]
				ATPase, H+ transporting,	
				lysosomal (vacuolar proton	ATPase, H+ transporting, lysosomal
1000	22661	Al169265	s,z	pump), subun	(vacuolar proton pump), subunit 1
1001	13239	Al169278	g,j,l,y,z		ESTs
1002	24162	Al169279	m		ESTs
					ESTs, Highly similar to Y069_HUMAN
					HYPOTHETICAL PROTEIN KIAA0069
1003	16879	AI169284	0		[H.sapiens]
1001	0.4045		L_		ESTs, Highly similar to H33_HUMAN
1004	24213	AI169289	p		HISTONE H3.3 [R.norvegicus]
1005	13240	AI169311	cc b		ESTs ESTs
1006	5931	AI169324	D		ESTS, Highly similar to CGI-117
1007	20891	A1160227	d		protein [H.sapiens]
1007	11979	AI169337 AI169365	cc		ESTs
1000	11313	M103303	 -	arachidonic acid	
1009	10947	AI169372	s	epoxygenase	arachidonic acid epoxygenase
1010	20697	AI169372	o,u	орохудопаво	ESTs
1011	8234	AI169517	z		ESTs
1012	18343	AI169648	0		ESTs
1012	10839	AI169655	l,m		ESTs
	+	1	t	 	ESTs, Weakly similar to hypothetical
	1		1		LO13, Weakly Silling to Hypothetical

Sequence ID No. 1015 1016 1017 1018	22575 804 8213	GenBank Acc/ Ref. Seq ID : ::	Modal Gods	Gene Name	
1015 1016 1017	22575 804			See the resident of the state o	Unigene Cluster Title
1017			r		ESTs, Moderately similar to T47184 hypothetical protein DKFZp434F1526.1 [H.sapiens]
	8213	AI169756	СС		ESTs, Highly similar to GENE 33 POLYPEPTIDE [R.norvegicus]
	2040	AI169883	p i bb		ferritin light chain 1
	3916	AI169947	i,bb		ESTs ESTs
1019 1020	3733 14179	AI170053 AI170224	u,General cc	1	ESTs
1021	11406	AI170224	r		ESTs, Moderately similar to class II cytokine receptor 4 [M.musculus]
1021	3547	AI170263	General		ESTs, Weakly similar to ZNT1 RAT ZINC TRANSPORTER 1 [R.norvegicus]
1023	11524	Al170340	j,y,z		ESTs, Weakly similar to CL36 RAT LIM DOMAIN PROTEIN CLP-36 [R.norvegicus]
1024	2729	AI170363	e,i		ESTs
1025	18811	AI170525	i		ESTs
1026	22524	AI170542	h		ESTs ESTs, Highly similar to CGI-10 protein
1027	24048	AI170570	a,g		[H.sapiens] ESTs, Moderately similar to AF161588
1028	5968	AI170692	y,aa		1 GABA-A receptor-associated protein [R.norvegicus]
1029	9757	AI170693	b		ESTs
1030	18905	Al170770	e,s		ESTs, Highly similar to NADH- ubiquinone oxidoreductase NDUFS2 subunit [H.sapiens] ESTs, Moderately similar to
1031	16170	AI170894	i		adipophilin [H.sapiens] Hyaluronan mediated motility receptor
1032	7089	Al171185	С		(RHAMM)
1032	17591	Al171185	b		ESTs
			-		ESTs, Weakly similar to AIF-C1
1034	13285 4428	AI171361 AI171362	a	(ubiquinone) Fe-S protein 1 (75kD) (NADH-coenzyme Q	[R.norvegicus] ESTs, Moderately similar to NUAM_HUMAN NADH-UBIQUINONE OXIDOREDUCTASE 75 KD SUBUNIT PRECURSOR [H.sapiens]
1036	18126	AI171369	w		ESTs, Highly similar to S16788 probable reverse transcriptase - rat [R.norvegicus]
1037	23253	Al171448	o		ESTs, Moderately similar to 68MP MOUSE 6.8 KD MITOCHONDRIAL PROTEOLIPID [M.musculus]
1038	4584	Al171492	m, General		ESTs
1039	11158	AI171432	r,s		ESTs, Moderately similar to NADH:ubiquinone oxidoreductase B22 subunit [H.sapiens]
	15345	AI171542	1		ESTs
	21183	AI171676	k		ESTs
1042	8215 11437	AI171692 AI171794	i i	ferritin light chain 1	Rattus norvegicus kynurenine aminotransferase/glutamine transaminase K (Kat) gene, complete cds,ferritin light chain 1 ESTs
		Al171800	cc		ESTs
	23579	AI171802	٧		ESTs
1046	11708	Al171807	l,t		ESTs
1047	17204	Al171844	s,y,z	HMm:RIKEN cDNA	Rattus norvegicus F1-ATPase epsilon subunit mRNA, nuclear gene encoding mitochondrial protein, complete cds

TABLE 1: 8	SUMMARY				Atty. Docket No. 44921-5099WG Doc. No. 1793997.1
Sequince ID No.	[clentifier	Conbenk Acel Ref. Seq ID	Model Model	Gene Name	Unigene Cluster Mile
1048	4420	Al171916	m		ESTs
					ESTs, Highly similar to T08675 hypothetical protein DKFZp564F0522.1 [H.sapiens]
1049	3266	AI171948	l,m t		ESTs
1050 1051	19012 11205	AI172056 AI172057	1.		ESTs
		<u> </u>	a,q,bb		ESTs, Weakly similar to T33238 hypothetical protein T10H9.3 -
1052	6057	AI172102	b		Caenorhabditis elegans [C.elegans]
1053	19128	AI172103	m		Rat mRNA for 5E5 antigen, complete cds
1054 1055	6630	Al172107 Al172184	z n	 	ESTs
1056	11968	Al172208	bb		ESTS, Weakly similar to FETA RAT ALPHA-FETOPROTEIN PRECURSOR [R.norvegicus]
1057	6974	AI172263	I,m		ESTs
1057	23313	AI172203 AI172271	d		ESTs
1059	2140	Al172272	General		ESTs, Moderately similar to A53004 transcription elongation factor S-II - rat [R.norvegicus] ESTs, Weakly similar to S43056
1060	15382	AI172302	I,p, General		hypothetical protein - mouse [M.musculus]
1061	18689	AI172329	l		ESTs
1062	17887	Al172414	0		Rattus norvegicus apoptosis- regulating basic protein mRNA, complete cds
1062	17007	A1172414			ESTs, Highly similar to A44437 regenerating liver inhibitory factor
1063	3042	AI172447	General	HMm:isocitrate	RL/IF-1 - rat [R.norvegicus] ESTs, Weakly similar to IDHC RAT
1001	17004	11170101		dehydrogenase 2 (NADP+),	ISOCITRATE DEHYDROGENASE
1064	17291	AI172491	bb	mitochondrial	[R.norvegicus]
1065	26222	AI172506	р		ESTs
1066 1067	13095 8795	AI172595 AI172618	General		ESTS
1068	6454	AI175342	j,l,m,y		ESTs, Weakly similar to T31067 BIR repeat containing ubiquitin-conjugating enzyme BRUCE - mouse [M.musculus]
1070	4445	Al175466	x		ESTs, Highly similar to RRAS MOUSE RAS-RELATED PROTEIN R-RAS [M.musculus]
					ESTs, Highly similar to NHPX RAT NHP2/RS6 FAMILY PROTEIN
1071	3418	AI175475	m		YEL026W HOMOLOG [R.norvegicus] ESTs, Moderately similar to AF145050 1 translation elongation factor 1-delta
1072	18507	AI175551	bb		subunit [R.norvegicus]
1073	10217	AI175628	w		ESTs
1074	7262	AI175833	j,m,x		ESTs
1075	19004	Al175875	r		ESTs
1076	22352	Al175959	I,General		ESTs
1077	7022	AI176041	h,n		ESTs, Highly similar to pirin [H.sapiens] ESTs, Weakly similar to tazarotene-
1078	21467	AI176061	lt		induced gene 2 [H.sapiens]
1079	18581	AI176160	General		ESTs
1080	14159	AI176169	g		ESTs
1081	21742	AI176172	w		ESTs
					ESTs, Highly similar to P55-C-FOS PROTO-ONCOGENE PROTEIN
1082	10182	AI176185	Canaral		[R.norvegicus]
1083	22765	AI176265	General	L	ESTs

TABLE 1: S			THE PERSON NAMED AND POST OF		Aity, Docket No. 44921-5039Wo Doc. No. 1793397.1
ID (yo: Sequence	ldeniiier-	ConBenk Ace Ref. Seg ID	Model Gode	Gene Name	Unigene Guster fille
					ESTs, Weakly similar to GSHH RAT PHOSPHOLIPID HYDROPEROXIDE GLUTATHIONE PEROXIDASE
1084	6905	AI176275	а		[R.norvegicus]
1085	12999	AI176276	cc		ACETYLHEXOSAMINE PYROPHOSPHORYLASE [H.sapiens]
					ESTs, Highly similar to SMD2_HUMAN SMALL NUCLEAR RIBONUCLEOPROTEIN SM D2
1086	16438	AI176294	е		[H.sapiens]
1087	21130	AI176298	у		ESTs
1088	3014	AI176362	е		ESTs
1089	15015	AI176363	r		ESTs
1090	19006	AI176393	×		ESTs ESTs, Moderately similar to QPS1
1091	20001	AI176396	0		[H.sapiens]
1092	12174	AI176435	j,m		ESTs
1093	15191	AI176456	b,o,t,v,cc		Rat metallothionein-2 and metallothionein-1 genes, complete cds
1094	24236	AI176473	d,General		ESTs
1005	10510	A1470540			ESTs, Moderately similar to HS9B RAT HEAT SHOCK PROTEIN HSP
1095 1096	16518 2161	AI176546 AI176592	General		90-BETA [R.norvegicus]
1090	2101	A1170592	General		ESTs, Weakly similar to S63220 probable membrane protein YNL247w
					- yeast (Saccharomyces cerevisiae)
1097	12436	AI176610	General I,v,		[S.cerevisiae]
1098	2536	AI176616	General		ESTs
1099 1100	18525 23449	AI176792 AI176828	g g		ESTs ESTs
1101	23299	AI176839	General		ESTs
1102	3580	AI176848	е	<u> </u>	ESTs
1103	22103	Al176849	d,General		ESTs
1104	16036	AI176855	ļt		ESTs ESTs, Highly similar to
					phosphomannomutase Sec53p
1105	15588	AI176916	General		homolog [M.musculus]
1106	16917	AI176951	t		ESTs
1107	16124	AI176963	сс		Rattus norvegicus transcription factor MRG1 mRNA, complete cds
1108	15146	AI176969	b,General		ESTs
1109	5786	AI177058	f		ESTs, Weakly similar to PSE-binding factor PTF delta subunit [H.sapiens]
1110	2852	AI177059	С		ESTs ESTs, Highly similar to AF139894 1
1112	3156	AI177092	9		RNA-binding protein alpha-CP1 [M.musculus]
				HMm:adenine	Rat adenine phosphoribosyltransferase (APRT)
1113	14384	AI177096	а	phosphoribosyl transferase	gene, complete cds ESTs, Weakly similar to C1QB RAT
1114	12240	A1177110	General		COMPLEMENT C1Q SUBCOMPONENT, B CHAIN PRECURSOR [R.norvegicus]
1114	13310	Al177119	General		ESTs, Highly similar to CGI-10 protein
1115	24049	AI177341	g,p,s,u		[H.sapiens]
1116	15964	AI177360	o,General	<u> </u>	ESTs
1117	14989	AI177366	u	Integrin, beta 1	Integrin, beta 1
1118	7975	AI177374	aa	L	ESTs

TABLE 1:	SUMMARY				Aiiy, Dockei No. 44921-5039W0 Doc. No. 1793397.
Sequence		GenBank Acc/	Model		
Sequence	ldentifer	Combank Acc/ Ref. Seq ID.	Code Model	Cene Name	Unitene Cluster Title
	in transmer	Man Osque			Rattus norvegicus substrate binding
					subunit of type II 5'-deiodinase D2p29
1119	3006	AI177395	k		mRNA, complete cds
					Rattus norvegicus mRNA for hnRNP
1120 1121	9521	AI177683 AI177706	<u>г</u>		protein, partial ESTs
1121	- 3321	AITITIO	 	···	2013
1122	14425	AI177755	g,General		ESTs
1123	10611	AI177790	j,m		ESTs
					ESTs, Moderately similar to S27962 modulator recognition factor 1
1124	5356	AI177813	СС		[H.sapiens]
		7.11.10.10			ESTs, Highly similar to SAS_HUMAN
					SARCOMA AMPLIFIED SEQUENC
1125	11791	AI177843	General		[H.sapiens]
1126	14484	AI177867	General		ESTs, Weakly similar to putative eps protein [R.norvegicus]
1120	17707	71111001	Jonetai		ESTs, Weakly similar to DRAL
1127	5780	Al177869	General		[R.norvegicus]
					ESTs, Highly similar to TGIF MOUSE
1100	19184	A147900E	Conoral		5'-TG-3' INTERACTING FACTOR [M.musculus]
1128	19164	Al178025	General		ESTs, Moderately similar to C17orf1
1129	6059	AI178245	c,General		protein [H.sapiens]
		1			ESTs, Weakly similar to hypothetical
1130	23248	AI178267	У		protein [H.sapiens]
					ESTs, Weakly similar to
				1	YAE6_YEAST HYPOTHETICAL 13.4 KD PROTEIN IN ACS1-GCV3
1131	4073	AI178272	0		INTERGENIC REGION [S.cerevisiae]
1132	7838	Al178291	е		ESTs
1133	18996	AI178326	У		ESTs
					ESTs, Highly similar to I49523 Mouse primary response gene B94 mRNA,
1134	22488	AI178392	ь		3'end - mouse [M.musculus]
1135	18800	AI178504	n,p,aa		ESTs
1136	22197	AI178527	g,General		ESTs ESTs, Highly similar to MCM3
					MOUSE DNA REPLICATION
					LICENSING FACTOR MCM3
1137	3401	AI178684	bb		[M.musculus]
1138	17713	AI178700	m		ESTs
1139	14874	AI178735 AI178746	e v,General	<u> </u>	ESTs ESTs
1140	23567	7/11/0/40	, Jeneral	-	Rattus norvegicus alpha-globin (GloA)
1141	18907	AI178971	c		gene, complete cds
1142	20991	AI178979	ļi		ESTs
4440	5007	A1470000	L.,		ESTs, Moderately similar to Vanin-1
1143	5887	Al179099	q,t b,e,		[M.musculus]
1144	8477	AI179167	General		ESTs
1145	3348	AI179288	u,v		ESTs
1146	13608	AI179314	е		ESTs
1147	8849	AI179315	g.p		ESTs
1148	13611	AI179378	v,General		Rattus norvegicus mRNA for prostasin precursor, complete cds
1149	15438	AI179378	m,x	collagen type V, alpha 2	collagen type V, alpha 2
	· · · · · · ·			<u> </u>	ESTs, Moderately similar to RB17
			e,t,		MOUSE RAS-RELATED PROTEIN
1150	13614	Al179407	General		RAB-17 [M.musculus]
1151	15042	Al179422	b,General		ESTs
1152	2768	AI179481	i,General		ESTs
1153	24041	Al179580	b,i	·····	ESTs

TABLE 1:	SUMMARY .				ÁMy. Doeket No. 44921-5079W Doe. No. 1793897
Seguañae Id No.	ldeniffier.	Genbank (Acc) Ref. Seq ID	Model Gode	Cono Namo	Unigene Cluster Title
1154	19822	Al179599	o,General		R.norvegicus mRNA for ras-related GTPase Rab29
1155	23270	Al179601	q,General		ESTs
1156	5901	AI179605	е		ESTs
1157	16081	AI179610 AI179717	g,i,p	Heme oxygenase	Heme oxygenase
1158	14564	AI179717 AI179750	k General		ESTs ESTs
1159 1160	7918 6647	AI179795			ESTs
1100	0047	A1179795	g	hypothetical protein	2318
1161	9097	Al179875	o,General	LOC56728	hypothetical protein LOC56728 ESTs, Highly similar to GAP
1162	23989	Al179953	a		JUNCTION BETA-2 PROTEIN [R.norvegicus]
1163	12899	Al179967	b		ESTs
1164	1687	AI179971	c	Hemoglobin, alpha 1	Hemoglobin, alpha 1
1165	22569	AI179979	General		ESTs
1166	23514	Al179986	o,General	HHs:phosphoserine phosphatase	ESTs, Highly similar to L-3- phosphoserine phosphatase [H.sapiens]
			1 _		
1167	15892	AI179988	c,General		ESTs
1168	12402	AI180004	g		ESTs, Highly similar to Unknown [H.sapiens]
1169	5443	AI180165	General		ESTs, Moderately similar to testis specific DNAj-homolog [M.musculus] ESTs, Highly similar to A Chain A,
1170 1171	5481 24028	Al180170 Al180239	General		The Crystal Structure Of Human Eukaryotic Release Factor Erf1-Mechanism Of Stop Codon Recognition And Peptidyl-Trna Hydrolysis [H.sapiens]
1171	24020	A1100233	'		ESTs, Moderately similar to JC4978
1172	17089	AI180281	g		oxidative stress protein A170 - mous [M.musculus]
					ESTs, Moderately similar to Y273_HUMAN HYPOTHETICAL PROTEIN KIAA0273 [H.sapiens]
1173	3701	A1180306	aa		ESTs [H.sapiens]
1174	3352	Al180334	m		ESTS, Highly similar to AF114169 1 nucleotide-binding protein short form
1175	24368	Al180392	I,m		[M.musculus] ESTs, Moderately similar to SPA1
1176	14337	Al180414	c		MOUSE GTPASE-ACTIVATING PROTEIN SPA-1 [M.musculus]
<u></u>					Rattus norvegicus chemokine CX3C
1177	19080	Al227647	j,y,z		mRNA, complete cds
1178	22838	Al227667	aa		ESTs
1179	6765	Al227761	i,General		ESTs, Highly similar to T00367 hypothetical protein KIAA0665 [H.sapiens]
11/3	0705	/11/4/101	i, Coneiai		ESTs, Weakly similar to AF187065 1 p75NTR-associated cell death
1180	24054	Al227867	General		executor [R.norvegicus]
1181	7324	Al227885	i		ESTs
1182	23898	Al227987	d	Peptidylglycine alpha-	ESTs Peptidylglycine alpha-amidating
1183	1651	AI228068	n,w	amidating monooxygenase	monooxygenase
1184	14237	AI228128	е		EST
					ESTs, Weakly similar to C21I_HUMAN PUTATIVE PROTEIN
1185	14242	AI228197	General		C21ORF18 [H.sapiens]
1186	16913	AI228236	0		ESTs
1187	22915	AI228299	r		ESTs, Highly similar to p97 homologous protein [H.sapiens]

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TABÜE 18 S	OMMARY :				Atty, Docket No. 44921-503 Doc. No. 1793		
Sequence In Ma		Conenk Acc	Model .				
ID Kor	ldeniiier?	Ref. Seq ID	Code 🕌	Cone Name	Unigene Cluster Tille		
1188	8917	AI228301	General		ESTs		
1189	15879	AI228313	r,General		ESTs Wealth similarts		
					ESTs, Weakly similar to AFG1 YEAST AFG1 PROTEIN		
1190	13727	AI228326	o,General	· .	[S.cerevisiae]		
1191	6102	AI228335	General		ESTs		
					ESTs, Weakly similar to S70642		
1192	13730	AI228356	а		ubiquitin ligase Nedd4 - rat [R.norvegicus]		
1193	13745	AI228494	b,cc		[EST		
1194	4217	Al228587	s		ESTS, Weakly similar to M172_HUMAN MEMBRANE COMPONENT, CHROMOSOME SURFACE MARKER 2 [H.sapien		
-					ESTs, Weakly similar to T16757 hypothetical protein R144.3 -		
1195	16053	AI228596	сс		Caenorhabditis elegans [C.elegar		
1196	3557	Al228672	е		ESTs		
1197	11605	AI228682	e		ESTs		
1198	13203	Al228728	r		ESTs ESTs, Highly similar to protein inhibitor of activated STAT protein		
1199	13771	Al228848	g		PIAS1 [H.sapiens]		
1200	5918	Al229036	ŗ		ESTs		
1201	8235	Al229154	k	Vasiale constitution	ESTs		
				Vesicle-associated membrane protein	Vesicle-associated membrane pro		
1202	16203	Al229196	r	(synaptobrevin 2)	(synaptobrevin 2)		
1203 1204	13826 13144	Al229304 Al229320	g g		ESTs ESTs		
1204	4640	AI229320 AI229404	x.aa		ESTs .		
		101	.,		ESTs, Moderately similar to MKK MOUSE MAP KINASE-ACTIVATI		
1206	23563	Al229421	1		PROTEIN KINASE 2 [M.musculu: ESTs, Moderately similar to NAD		
					ubiquinone oxidoreductase PDSV		
1207	15426	AI229497	S		subunit [H.sapiens]		
1208	15193	Al229508	bb		ESTs ESTs, Highly similar to KITH RAT		
					THYMIDINE KINASE, CYTOSOL		
1209	19243	AI229638	X		[R.norvegicus] ESTs		
1210	23078	Al229647	р	HHs:NADH dehydrogenase	EOIS		
				(ubiquinone) Fe-S protein 3 (30kD) (NADH-coenzyme Q	ESTs, Highly similar to NADH:ubiquinone oxidoreductase		
1211	3099	AI229680	0	reductase)	NDUFS3 subunit [H.sapiens] Sprague-Dawley D-beta- hydroxybutyrate dehydrogenase		
1212	19508	Al229698	bb		mRNA, complete cds Rattus norvegicus mRNA for clas		
1213	13977	Al229707	x		beta-tubulin, complete cds ESTs, Moderately similar to		
					NADC_HUMAN NICOTINATE- NUCLEOTIDE		
1214	23983	Al229708	v		PYROPHOSPHORYLASE [H.sap		
1215	2688	Al229793	е		ESTs Wookly similar to KIAAOSE		
1216	13874	AI229832	g		ESTs, Weakly similar to KIAA085 protein [H.sapiens] ESTs, Weakly similar to MOT2 R.		
1017	12507	A1220070	Conoral		MONOCARBOXYLATE		
1217 1218	12587 20591	AI229979 AI229993	General I,m		TRANSPORTER 2 [R.norvegicus		
1218	24042	AI239993	a,b,d, General		ESTs		
	1		T		Rattus norvegicus mRNA for volta		
1220	13880	AI230042	u	I	gated ca channel, complete cds		

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TABLE 1: 8					. Ally. Docket No. 44921-5039W0
					Doc. No. <u>1793397.1</u>
Sequence		GenBank Acc	Model		esteric afficier de Very XII est
ID No:	(Identifier	Ref. Seq ID	Code	Gene Name	Unigene Gluster Tille
					ESTs, Highly similar to NIMM MOUSE
				HMm:NADH dehydrogenase	NADH-UBIQUINONE
1221	17672	AI230074	d	(ubiquinone) 1 alpha subcomplex, 1	OXIDOREDUCTASE MWFE SUBUNIT [M.musculus]
1221	17072	A1230074	-	Subcomplex, 1	Rattus norvegicus hfb2 mRNA,
1222	3652	AI230113	General		complete cds
					ESTs, Weakly similar to HS9B RAT
4000	10050	A1000404			HEAT SHOCK PROTEIN HSP 90-
1223	18650	Al230121	aa		BETA [R.norvegicus] ESTs, Moderately similar to
}					CHD3 HUMAN CHROMODOMAIN
					HELICASE-DNA-BINDING PROTEIN
1224	13025	AI230173	С		3 [H.sapiens]
1225	4280	AI230247	Z Conservation	selenoprotein P, plasma, 1	selenoprotein P, plasma, 1
1226	18528	AI230284	General		ESTs ESTs, Moderately similar to T46458
					hypothetical protein DKFZp434M102.1
1227	7084	AI230362	р		[H.sapiens]
1228	20895	Al230549	b,n		ESTs
1229	12961	AI230554	General		ESTs
1,000					Rattus norvegicus mRNA for galectin-
1230	15636 4121	Al230616 Al230647	j,m		2 related protein, complete cds ESTs
1231	4121	A1230047	1,,,,,		ESTs, Highly similar to HN1
1232	14388	AI230702	General		[M.musculus]
1233	18529	Al230716	x,General		ESTs
					Rattus norvegicus phosphoinositide
1,004	10010				phosphatase SAC1 mRNA, complete
1234 1235	13618 8304	Al230724 Al230746	General		cds ESTs
1236	4731	Al230773	e		ESTs
		72007.10			ESTs, Moderately similar to
					CDN3_HUMAN CYCLIN-
1	1		١.		DEPENDENT KINASE INHIBITOR 3
1237	14430	AI230798	c,k,x		[H.saplens] ESTs, Highly similar to AF102850 1
				HHs:Alg5, S. cerevisiae,	dolichyl-phosphate beta-
1238	16627	AI230822	bb	homolog of	glucosyltransferase [H.sapiens]
					Rattus norvegicus mRNA for brain
1239	3125	Al231028	General		4.1(S), complete cds
1240	633	Al231127	k		ESTs
					ESTs, Highly similar to RL2B_HUMAN 60S RIBOSOMAL PROTEIN L23A
1241	20846	AI231140	р		[R.norvegicus]
1242	6743	Al231219	d		ESTs
1244	26292	Al231391	k		
1245	12343	AI231433	w		ESTs
1246 1247	7337	AI231465	aa General		ESTs ESTs
1241	16321	AI231506	General		ESTs, Highly similar to Z183_HUMAN
					ZINC FINGER PROTEIN 183
1248	8004	AI231532	j,1		[H.sapiens]
					ESTs, Moderately similar to BAG-
					family molecular chaperone regulator-
1249	15171 6193	Al231792 Al231797	g		3 [H.sapiens] ESTs
1230	0193	A1231797			ESTs, Moderately similar to tumor
1252	14227	AI231999	u		protein D53 [M.musculus]
					Rattus norvegicus translation
					elongation factor 1-delta subunit
1253	24501	AI232006	w,y,bb		mRNA, partial cds
1254	3434	A1232014	g,q,z,cc,		ESTs
1254	13434	AI232014	General		ESTs, Highly similar to Human
					Translation Initiation Factor Eif1, Nmr,
1255	19094	Al232021	n,General		29 Structures [H.sapiens]

TABLE 1: S	UMMARY				Mily. Docket No. 44921-5066110
				1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	Doc. No. 1798397.1
Sequence ID No.	lienilier	Consont Acci Ref. Seq ID	Model Code	Cene Name	Unione Circler Title
1256	14020	Al232076	u		ESTs
1257	6726	Al232157	d		ESTs
1258	11549	Al232174	l,m		ESTs
1259	23125	Al232266	j,s		ESTs ESTs, Moderately similar to JC4914
1260	2085	Al232270	bb		anti-sigma cross-reacting protein homolog I beta precursor [H.sapiens]
1261	2913	AI232272	О		ESTs, Weakly similar to T25417 hypothetical protein T28D6.9 - Caenorhabditis elegans [C.elegans]
1262	14304	AI232281	g		ESTs, Weakly similar to KIAA0971 protein [H.sapiens]
1263	15955	AI232294	u,bb, General		ESTs
1264	15122	AI232303	у		ESTs, Weakly similar to Sid1669p [M.musculus]
1265	4716	AI232313	у	purinergic receptor P2X, ligand-gated ion channel 4	purinergic receptor P2X, ligand-gated ion channel 4
1266	15246	AI232332	t,u o	Stromal cell-derived factor 1	ESTs Stromal cell-derived factor 1
1267	16172	Al232340 Al232341	d	Stromal cell-derived factor 1	ESTS, Weakly similar to YQ42_CAEEL HYPOTHETICAL 40.0 KD PROTEIN C13B9.2 IN CHROMOSOME III [C.elegans]
1269	11411	Al232346	h		ESTs
1270	19287	Al232379	f	Platelet-derived growth factor receptor alpha	Platelet-derived growth factor receptor alpha
1271	5601	Al232461	n,General		ESTs, Weakly similar to FMO1 RAT DIMETHYLANILINE MONOOXYGENASE [R.norvegicus] ESTs, Weakly similar to PIR1
1272	14051	AI232489	I,m		[H.sapiens] ESTs, Moderately similar to A27340
1273	5572	AI232490	i,t		complement C7 precursor [H.sapiens]
1274	11157	Al232494	СС		ESTs
1075	0700	11000504			ESTs, Weakly similar to DnaJ
1275 1276	8709 20350	Al232534 Al232552	j,v,y		homolog 2 [R.norvegicus]
1277	14069	AI232631	е		ESTs
1278	4440	AI232643	w		ESTs ESTs, Weakly similar to putative peroxisomal 2.4-dienoyl-CoA
1279	17695	Al232784	e		reductase [R.norvegicus]
1280	15796	Al232874	V		ESTs
1281	12467	Al232924	General		ESTs
1282	12873	AI232984	ļi		ESTs
1283	5355	AI233031	r		ESTs ESTs, Moderately similar to MHC
1284	18794	Al233121	b,g,		class I [M.musculus] ESTs, Weakly similar to nuclear RNA
1285	3823	Al233147	General c,k,		helicase [R.norvegicus]
1286	11967	AI233155	General		ESTs
1287	11561	AI233182	d		ESTS, Highly similar to PM1_HUMAN PROTEIN PM [H.sapiens]
1288	3471	AI233183	9		ESTs, Weakly similar to T15919 hypothetical protein EEED8.9 -
1289	21948	Al233191	i		Caenorhabditis elegans [C.elegans]
1290	13598	Al233194	g,p,y		ESTs
1291	15552	Al233195	у		ESTs, Highly similar to Bodenin [M.musculus]
					Rattus norvegicus epidermal growth factor receptor related protein (Errp)
1292	17907	AI233224	bb		mRNA, complete cds
1293	14111	Al233269	cc		ESTs

TABLE 9: 8	BUMMARY				:- Aliy. Dockel No. 44921-5089000 Doc. No. 1798897.
Sequence ID'No.		ConBank Ace Ref. Seq ID	Model		
ID No.	ldentifier	Rei-Seg ID	Code	Gene Name	Unigene Cheter Tille
					ESTs, Weakly similar to T24956
		, V, V, V, V			hypothetical protein T16G1.10 -
1294	12894	AI233365	d		Caenorhabditis elegans [C.elegans]
					ESTs, Weakly similar to S44853
					K12H4.3 protein - Caenorhabditis
1295	7161	AI233407	General		elegans [C.elegans]
1296	15906	AI233425	q		ESTs
1297	14120	AI233433	d		ESTs
1298	14095	AI233468	a,d		ESTs
1230	14030	711200400	14,4		ESTs, Weakly similar to I38079 OXA1
1299	3075	AI233494	22		homolog [H.sapiens]
			u,aa Conorol		ESTs
1300	6046	AI233530	General		
					PSD8_HUMAN 26S PROTEASOME
					REGULATORY SUBUNIT S14
1301	18900	AI233570	General		[H.sapiens]
					ESTs, Moderately similar to
			į.	HHs:arginyl-tRNA	SYR HUMAN ARGINYL-TRNA
1302	7888	AI233583	General	synthetase	SYNTHETASE [H.sapiens]
1303	16709	AI233602	General	Adenosin kinase	Adenosin kinase
1303	10109	\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	Jeneral	AGCIIOSIII KIIIdSE	ESTs, Highly similar to
			I		
			İ		P2CD_MOUSE PROTEIN
	İ				PHOSPHATASE 2C DELTA
					ISOFORM (PP2C-DELTA) (P53-
	1		l .		INDUCED PROTEIN PHOSPHATASE
					1) (PROTEIN PHOSPHATASE
					MAGNESIUM-DEPENDENT 1
1204	5163	AI233712	v		DELTA) [M.musculus]
1304	5163	A12337 12	У		
			Ì		ESTs, Moderately similar to ERHUAH
					coatomer complex alpha chain
1305	7243	AI233717	General		homolog [H.sapiens]
					ESTs, Highly similar to
					PSD5 HUMAN 26S PROTEASOME
1306	3816	AI233729	g		SUBUNIT S5B [H.sapiens]
1000	10010	7.1.2007.20	13	· · · · · · · · · · · · · · · · · · ·	ESTs, Weakly similar to ALDR RAT
			ا ا		ALDOSE REDUCTASE
	1,,,,,,		d,h,		
1307	13023	AI233740	General		[R.norvegicus]
1308	14871	AI233743	g		ESTs
					ESTs, Highly similar to Gene product
					with similarity to KIAA0154
1309	7469	AI233767	cc		[H.sapiens]
1310	7804	Al233771	b		ESTs
		1	-		ESTs, Weakly similar to T24413
					hypothetical protein T04A11.2 -
1011	40500	41000770	1_		
1311	13563	A1233773	e	ļ	Caenorhabditis elegans [C.elegans]
1312	2154	AI233818	k,cc		ESTs
1313	16616	AI234079	h		ESTs
		·	a,d,		
1314	13393	Al234100	General	cysteine rich protein	cysteine rich protein
1315	7071	AI234162	r		ESTs
1316	14677	Al234620	General		EST
	17011	7.11207020	Jonesia		ESTs, Weakly similar to transcription
1310		i	l		
	4442	A100460C		1	factor C1 [M.musculus]
1317	4443	AI234629	m		IFOT-
1317 1318	22453	Al234678	b		ESTs
1317 1318 1319	22453 23964	Al234678 Al234748			ESTs
1317 1318 1319	22453	Al234678	b		
1317 1318 1319	22453 23964	Al234678 Al234748	b		ESTs
1317 1318 1319 1320	22453 23964 19581	Al234678 Al234748 Al234753	b t,General f	DEXRAS1 (Dexras1)	ESTs
1317 1318 1319 1320	22453 23964	Al234678 Al234748	b t,General f	DEXRAS1 (Dexras1)	ESTs EST DEXRAS1 (Dexras1)
1317 1318 1319 1320	22453 23964 19581 22152	A1234678 A1234748 A1234753 A1234822	t,General f o,General	DEXRAS1 (Dexras1)	ESTs EST DEXRAS1 (Dexras1) ESTs, Weakly similar to S12207
1317 1318 1319 1320	22453 23964 19581	Al234678 Al234748 Al234753	b t,General f		ESTs EST DEXRAS1 (Dexras1)
1317 1318 1319 1320	22453 23964 19581 22152	A1234678 A1234748 A1234753 A1234822	t,General f o,General	ATPase, H+ transporting,	ESTs EST DEXRAS1 (Dexras1) ESTs, Weakly similar to S12207 hypothetical protein [M.musculus]
1317 1318 1319 1320	22453 23964 19581 22152	A1234678 A1234748 A1234753 A1234822	t,General f o,General	ATPase, H+ transporting, lysosomal (vacuolar proton	ESTS EST DEXRAS1 (Dexras1) ESTs, Weakly similar to S12207 hypothetical protein [M.musculus] ATPase, H+ transporting, lysosomal
1317 1318 1319 1320 1321	22453 23964 19581 22152	A1234678 A1234748 A1234753 A1234822	t,General f o,General	ATPase, H+ transporting,	ESTs EST DEXRAS1 (Dexras1) ESTs, Weakly similar to S12207 hypothetical protein [M.musculus] ATPase, H+ transporting, lysosomal (vacuolar proton pump), subunit 1
1317 1318 1319 1320 1321	22453 23964 19581 22152 18942	A1234678 A1234748 A1234753 A1234822 A1234865	b t,General f o,General d	ATPase, H+ transporting, lysosomal (vacuolar proton	ESTS EST DEXRAS1 (Dexras1) ESTs, Weakly similar to S12207 hypothetical protein [M.musculus] ATPase, H+ transporting, lysosomal (vacuolar proton pump), subunit 1
1317 1318 1319 1320 1321 1322	22453 23964 19581 22152 18942	A1234678 A1234748 A1234753 A1234822 A1234865	b t,General f o,General d	ATPase, H+ transporting, lysosomal (vacuolar proton	ESTs EST DEXRAS1 (Dexras1) ESTs, Weakly similar to S12207 hypothetical protein [M.musculus] ATPase, H+ transporting, lysosomal (vacuolar proton pump), subunit 1 ESTs, Highly similar to CB80_HUMAN
1317 1318 1319 1320 1321	22453 23964 19581 22152 18942	A1234678 A1234748 A1234753 A1234822 A1234865	b t,General f o,General d	ATPase, H+ transporting, lysosomal (vacuolar proton pump), subun	ESTS EST DEXRAS1 (Dexras1) ESTs, Weakly similar to S12207 hypothetical protein [M.musculus] ATPase, H+ transporting, lysosomal (vacuolar proton pump), subunit 1

TABLE 1: 8	BUMMYARY				Anny, Docket No. 44921-503900 Doc. No. 1793397.
Sequence ID No.	ldentifier	Cenibank Acel Ref. Seq ID	Model Gode	©ene Name	Uniçin Civster Titlə
			117753		ESTs, Highly similar to ABF2_HUMA ATP-BINDING CASSETTE, SUB- FAMILY F, MEMBER 2 (IRON INHIBITED ABC TRANSPORTER 2)
1326	14906	Al235192	9		[H.sapiens]
1327	14718	Al235210	е		ESTs
1328	15004	AI235224	b,General		Rattus norvegicus tissue inhibitor of metalloproteinase-1 (TIMP1), mRNA, complete cds
1329	6632	AI235277	V		ESTs
1330	14722	AI235284	x,z		ESTs, Weakly similar to single-pass transmembrane protein [R.norvegicus Rat mRNA for preprocathepsin D (EC
1331	1462	AI235585	u,General		[3.4.23.5]
1332	21061	AI235631	I.m		ESTs
		<u> </u>	† · · · · · ·		
1333	14665	AI235646	m	MAD homolog 4 (Drosophila)	MAD homolog 4 (Drosophila) ESTs, Moderately similar to pescadill
1334	19940	AI235689	General		[H.sapiens]
1335	5698	Al235692	u		ESTs
1336	23745	AI235732	k		ESTs, Highly similar to NID2 MOUSE NIDOGEN-2 PRECURSOR [M.musculus]
1330	23743	A1233732	 		ESTs, Moderately similar to A56716
1337	11164	AI235739	General		aromatic ester hydrolase [H.sapiens]
1338	5212	AI235745	d		ESTs
					ESTs, Weakly similar to hypothetical
1339	14768	AI235912	h		protein [H.sapiens]
1340	14776	AI235950	m		ESTs
1341	3091	AI236027	n,General		ESTs
1342	14861	AI236045	r		ESTs EST
1343	14862	AI236048	е		ESTs, Highly similar to E25B protein
1344	16943	AI236097	р		[M.musculus] ESTs, Highly similar to JC7107
		1			development related unidentified 27k
1345	8336	AI236101	li .		protein - mouse [M.musculus]
1346	23230	Al236146	v		ESTs
1347	22855	AI236150	е		ESTs, Highly similar to JC7301 Dowr syndrome critical region protein 5 alpha [H.sapiens]
1348	14594	Al236152	li		ESTs
1349	18406	Al236168	r		ESTs
1350	15051	Al236332	General		ESTs, Highly similar to ATDA MOUS DIAMINE ACETYLTRANSFERASE [M.musculus]
	10209		hh		ESTs, Weakly similar to NHPX RAT NHP2/RS6 FAMILY PROTEIN YEL026W HOMOLOG [R.norvegicus
1351	19298	AI236338	bb		siah binding protein 1; FBP interaction
1352	10667	Al236366	b_	siah binding protein 1; FBP interacting repressor; pyrimidine tr	repressor; pyrimidine tract binding splicing factor; Ro ribonucleoprotein-binding protein 1
1353	10774	AI236397	f		ESTs
1354	9407	Al236402	aa		ESTs Rattus norvegicus retinol dehydrogenase type II mRNA,
1355	26335	Al236460	General		complete cds
1356	17950	Al236590	t,General		ESTs
1357	18259	AI236601	h,v		ESTs
1358	11445	Al236613	j,y		ESTs ESTs, Highly similar to SCF complex
1359	17248	AI236635	o,aa		protein Skp1 [M.musculus]
1360	16859	Al236753	t,General		ESTs

PCT/US02/16173

·				-90-	
TABLE 1:	SUMMARY				· Ally, Dockel No. 44921-5089WO
1.0					Dos. No. 1793397.1
Sequence			Model		Mark Colores
no Mer Sednevee	ldenillier.	Conbook Acc/ Ref. Seq ID	Goods (Mocion)	Cons Name	Unigene Cluster Tille
ID INO.	indentifier.	INGIA COGI ID	and the state of t	Catanenia	ESTs, Weakly similar to hT41
1361	5208	AI236754	g		[H.sapiens]
1362	24388	AI236772	e,General		ESTs
					ESTs,ESTs, Highly similar to HS9B
1363	15850	AI236795	n,v,w		RAT HEAT SHOCK PROTEIN HSP 90-BETA [R.norvegicus]
1364	14800	AI236856	w		ESTs
1366	11404	AI237002	m	spermidine synthase	spermidine synthase
4007	10454	A1027040	Conoral		ESTs, Highly similar to hepatitis B
1367	18151	AI237212	o,General	estrogen-responsive uterine	virus X interacting protein [H.sapiens]
1368	21653	AI237535	t,General	transcript	estrogen-responsive uterine transcript ESTs, Moderately similar to INIB RAT
					INTERFERON-INDUCIBLE PROTEIN
1369	11208	AI237586	z		[R.norvegicus]
					ESTs, Moderately similar to
			l.,		Y101_HUMAN HYPOTHETICAL
1370	21893	AI237713	i,k,aa		PROTEIN KIAA0101 [H.sapiens]
1371	14842	Al237724	ļr		ESTs ESTs, Moderately similar to MXI1 RAT
					MAX INTERACTING PROTEIN 1
1372	3467	AI237835	General	İ	[R.norvegicus]
1373	25840	Al638972	u		
			†		ESTs, Highly similar to G9A
1374	17108	AI639017	n		[M.musculus]
				mini chromosome	
				maintenance deficient 6 (S.	mini chromosome maintenance
1375 1376	16676 12400	Al639082 Al639107	c,k,x	cerevisiae)	deficient 6 (S. cerevisiae)
1377	19952	Al639108	q,v		ESTs
1379	25907	Al639167	0,W		ESTs
13,5		1	1-,		ESTs, Highly similar to T46480
					hypothetical protein
1381	18533	AI639231	n		DKFZp434L1850.1 [H.sapiens]
1382	18353	AI639233	t,aa	decorin	decorin
1384	15330	AI639285	General		ESTs EST
1385 1386	20026 25971	AI639354 AI639365	g r		[23]
1300	23971	A1039303	 		
1388	19152	AI639387	u,General		ESTs
					ESTs, Moderately similar to CAQC
					RAT CALSEQUESTRIN, CARDIAC
	1				MUSCLE ISOFORM PRECURSOR
1390	18338	Al639422	У		[R.norvegicus] [EST, Highly similar to A42772 mdm2
1392	20082	AI639488	i,m		protein - rat [R.norvegicus]
1032	120002	711000-100	,,	<u> </u>	ESTs, Weakly similar to T13607
			a,bb,		hypothetical protein EG:87B1.3 - fruit
1394	20056	AI639504	General	<u></u>	fly [D.melanogaster]
					ESTs, Highly similar to
					RPB8_HUMAN DNA-DIRECTED RNA
1205	4740	A1620540	_		POLYMERASES I, II, AND III 17.1 KD
1395	4713	Al639518	9	protein phosphatase 1,	POLYPEPTIDE [H.sapiens]
	1			regulatory (inhibitor) subunit	protein phosphatase 1, regulatory
1396	14332	AJ001044	bb	5	(inhibitor) subunit 5
1397	7602	AJ001929	k	reticulocalbin	reticulocalbin
					Rattus norvegicus mRNA for
1398	9867	AJ005424	u O '	lata dia 7	BMK1/ERK5 protein, partial
1400	16351	AJ011811	General	claudin 7 growth differentiation factor	claudin 7
1401	20116	AJ011969	I,General	15	growth differentiation factor 15
1.73	 -~	1,001,1909	,,conoral	· -	Rattus norvegicus mRNA for
1402	17635	AJ223355	v,w	l	mitochondrial dicarboxylate carrier
•				-	• • • • • • • • • • • • • • • • • • • •

TABLE 1: E					Ally, Dockel No. 44921-5039VVG Doc. No. 1793397.
Sequence ID No. i	ldendifier	GenBank Acc/ Ref. Seq ID	Model Code	Cene Name	Unigene Cluster Tille
1403	18686	D00729	q	dodecenoyl-Coenzyme A delta isomerase (3,2 trans- enoyl-Coenyme A	Rat mRNA for delta3, delta2-enoyl- CoA isomerase,dodecenoyl- Coenzyme A delta isomerase (3,2 trans-enoyl-Coenyme A isomerase)
1404	5049	D10655	n,w	dihydrolipoamide acetyltransferase	dihydrolipoamide acetyltransferase
1405	25257	D13623	<u> </u>		IESTs
1405 1406	15281 11434	D13623 D14014	h cc		ESTs
1407	1613	D14076	x		Rat mRNA for testicular dynamin, complete cds
1408	1728	D16479	q	HHs:hydroxyacyl-Coenzyme A dehydrogenase/3-ketoacyl- Coenzyme A thiolase/enoyl- Coenzyme A hydratase (trifunctional protein), beta subunit	
1409	3015	D16554	c,s,v,z		repetitive ubiquitins in tandem), complete cds
1410	472	D26111	d,s,bb		R.norvegicus mRNA for chloride channel (putative) 2313bp
1412	16233	D29960	j,l		Rattus norvegicus mRNA for alphaB crystallin-related protein, complete cds ESTs, Highly similar to PRC6 RAT
1413	9029	D30804	n		PROTEASOME SUBUNIT RC6-1 [R.norvegicus] Rattus norvegicus tyrosine
1414	1485	D38222	y,z		phosphatase-like protein IA-2a mRNA partial cds
1415	9135	D45247	s	proteasome beta type subunit 5	ESTs, Highly similar to PRCE RAT PROTEASOME EPSILON CHAIN PRECURSOR [R.norvegicus]
1416	16354	D50564	u	HHs:mercaptopyruvate sulfurtransferase	Rattus norvegicus mRNA for mercaptopyruvate sulfurtransferase, complete cds
1417	1884	D50695	l,m,bb		Rattus norvegicus mRNA for proteasomal ATPase (Tat-binding protein7), complete cds
				Solute carrier family 1 A1	Solute carrier family 1 A1 (brain
1419	826 826	D82928	General f	(brain glutamate transporter) HHs:CDP-diacylglycerol- inositol 3- phosphatidyltransferase (phosphatidylinositol synthase)	glutamate transporter) Rat mRNA for phosphatidylinositol synthase, complete cds
1420	25306 18867	D84485 D88250	u t		Rattus norvegicus mRNA for serine protease, complete cds
1423	22543	H31117	r,v, General		
1424	12360	H31456	w		ESTs
1425 1426	11358	H31489 H31610	h,j h		ESTs ESTs, Highly similar to mtprd [M.musculus]
1427	4360	H31813	bb, General		ESTs, Moderately similar to T14781 hypothetical protein DKFZp586B1621.1 [H.sapiens] ESTs, Moderately similar to COF1
	0242	H32169			RAT COFILIN, NON-MUSCLE ISOFORM [R.norvegicus]
1428 1429	9343 4386	H33093	h,w		EST [K:Norvegicus]

TABLE 1: S	UMMARY				Airy, Doctor No. 44921-5199740
				100000000000000000000000000000000000000	Doc. No. 1793397.1
Sequence ID No.	ldeniiier	Confenik Acc/ Ref. Seq ID	Model Code	Gene Name	Unigene Gluster Tille
	neemanes.	And the Control of th	b.date.e.millions		ESTs, Highly similar to IF39_HUMAN
					EUKARYOTIC TRANSLATION INITIATION FACTOR 3 SUBUNIT 9
1431	15374	H34186	1		[H.sapiens]
1432	17159	J00797	u,General	alpha-tubulin	alpha-tubulin
1433	16260	J01878	f		Rat brain-specific identifier sequence RNA, clone p1b224
1400	10200	001070		Branched chain alpha-	
1434	17284	J02827	bb	ketoacid dehydrogenase subunit E1 alpha	Branched chain alpha-ketoacid dehydrogenase subunit E1 alpha
1404	17204	302027		Subdiffic E 1 dipila	Rat glutathione S-transferase mRNA,
1435	15017	J03752	n	Thyroid hormone receptor,	complete cds
				beta (avian erythroblastic	Thyroid hormone receptor, beta (aviar
				leukemia viral (v-erb-a)	erythroblastic leukemia viral (v-erb-a)
1436	44	J03819	p,s e.r.	oncogene homolog 2) Glutathione-S-transferase,	oncogene homolog 2) Glutathione-S-transferase, mu type 2
1437	21014	J03914	General	mu type 2 (Yb2)	(Yb2)
				Steroid-5-alpha-reductase,	
				alpha polypeptide 1 (3-oxo-5 alpha-steroid delta 4-	Steroid-5-alpha-reductase, alpha polypeptide 1 (3-oxo-5 alpha-steroid
1438	20429	J05035	f	dehydrogenase alpha 1)	delta 4-dehydrogenase alpha 1)
1400	20423	000000	<u>'</u>	Glutamylcysteine gamma	Glutamylcysteine gamma synthetase
1439	1247	J05181	j,I,m,s,y,z	synthetase light chain	light chain
4440	10464	105540	n,u,	Inositol 1,4,5-triphosphate	Rat inositol-1,4,5-triphosphate receptor mRNA, complete cds
1440 1441	20149	J05510 K03243	General g	receptor type 1	receptor microx, complete cas
13-71	20143	100240	<u> </u>		Rat peroxisomal enoyl-CoA:
					hydrotase-3-hydroxyacyl-CoA
4440	47750		_		bifunctional enzyme mRNA, complete
1442 1443	17758 381	K03249 L00124	lq w	Elastase 2, pancreatic	cds Elastase 2, pancreatic
1444	2048	L00382	k,x	Liastase 2, parioreate	Ciastase 2, parioreate
1445	10500	L04619	s		
					Rattus norvegicus clone 15 polymeric immunoglobulin receptor mRNA,
1447	108	L14002	lp lt		3'UTR microsatellite repeats
1448	25366	L14003	l .		Rattus norvegicus clone 15 polymeric
					immunoglobulin receptor mRNA,
1449	109	L14004	с,р		3'UTR microsatellite repeats
1450 1451	20414 25369	L14323 L14937	General v	Phospholipase C-beta1	Phospholipase C-beta1
1451	25369	L14937	<u>y</u>	2',3'- Cyclic nucleotide 3'-	2',3'- Cyclic nucleotide 3'-
1452	16119	L16532	k	phosphodiesterase	phosphodiesterase
1453	25377	L25387	h		507 48 48 48 48 48 48 48 48 48 48 48 48 48
1453	12058	L25387	h		ESTs, Highly similar to A53047 6- phosphofructokinase [R.norvegicus]
. 100	12000		.,	Solute carrier family 1 A1	Solute carrier family 1 A1 (brain
1455	21146	L35558	General	(brain glutamate transporter)	glutamate transporter)
1456	106	L37203	w		Rattus norvegicus guanylyl cyclase (GC-D) mRNA, complete cds
					Rattus norvegicus serine protease
1458 1459	13682 6405	L38482 L38615	f,j,k,m,z p	Glutathione synthetase gene	gene, complete cds Glutathione synthetase gene
1461	15189	M11794	n,v	Cidatilone symmetase gene	Cistatrione symmetres gene
1462	17086	M13011	j		Rat c-ras-H-1 gene, complete cds
4404	04055	145404			Rat insulin-like growth factor-I mRNA,
1464 1465	21053 25405	M15481 M18330	j.l		3' end
1466	25415	M19648	a		
1468	14967	M22366	w		
				Propionyl Coenzyme A	
		i		carboxylase, alpha	

TABLE 1: S	ord Pos				Aijy. Docket No. 44921-51199W0 Doc. No. 1799397 1
Sequence ID No.	ldeniller	Conedia Ace	Model Gode	Cene Name	Unigene Cluster Tille
4.474	15048	M24542		HHs:ubiquinol-cytochrome c reductase, Rieske iron-sulfur polypeptide 1	Rat Rieske iron-sulfur protein mRNA, complete cds
1471 1472	20921	M29853	m m	рогурершие т	Rat cytochrome P-450 isozyme 5 (P450 IVB2) mRNA, complete cds
1473	1224	M31931	u	Cytochrome P450, an olfactory-specific steroid hydroxylase	Cytochrome P450, an olfactory- specific steroid hydroxylase
1474	15579	M33648	q	in yeroxy table	Rat mitochondrial 3-hydroxy-3- methylglutaryl-CoA synthase mRNA, complete cds
1474	15580	M33648	q		Rat mitochondrial 3-hydroxy-3- methylglutaryl-CoA synthase mRNA, complete cds
1475	17211	M34331	g,n,q,v		ESTs, Weakly similar to KRAB-zinc finger protein KZF-1 [R.norvegicus]
1476	20699	M35601	b,x,bb		Rat alpha-fibrinogen mRNA, 3' end
1476	20700	M35601	b,t,bb	 	Rat alpha-fibrinogen mRNA, 3' end
1477	9223	M36151	0		Rat mRNA for MHC class II antigen RT1.B-1 beta-chain,Rattus norvegicus MHC class II antigen RT1.B beta chain mRNA, partial cds Rat general mitochondrial matrix
1479	1585	M57728	j,m,y		processing protease (MPP) mRNA, 3'
1480	24844	M58040	С	transferrin receptor	transferrin receptor
1481	25057	M58495	h		
1482	457	M60666	d,General	Tropomyosin 1 (alpha)	Tropomyosin 1 (alpha) Rat cystatin S (CysS) gene, complete
1483	1223	M75281	f	P-glycoprotein/multidrug	cds
1484	5733	M81855	i,k,aa	resistance 1	P-glycoprotein/multidrug resistance 1 Rat beta-galactoside-alpha 2,6-
1485	4198	M83143	m		sialyltransferase mRNA Rat beta-galactoside-alpha 2,6-
1485	4199	M83143	m		sialyltransferase mRNA
1486	24651	M83678	k,x,z	RAB13	RAB13
1487	1430	M84648	General	Dopa decarboxylase (aromatic L-amino acid decarboxylase)	Dopa decarboxylase (aromatic L- amino acid decarboxylase)
1488	25467	M93297	С	ornithine aminotransferase	ornithine aminotransferase
1489	729	M95762	а,у		Rattus norvegicus GABA transporter GAT-2 mRNA, complete cds
				Rattus norvegicus Acetyl- CoA acyltransferase, 3-oxo acyl-CoA thiolase A, peroxisomal (Acaa), mRNA.	Acetyl-CoA acyltransferase, 3-oxo
1490	23698	NM_012489	q	Length = 1619 Rattus norvegicus Acetyl- CoA acyltransferase, 3-oxo acyl-CoA thiolase A, peroxisomal (Acaa), mRNA.	acyl-CoA thiolase A, peroxisomal Acetyl-CoA acyltransferase, 3-oxo
1490	23699	NM_012489	q	Length = 1619	acyl-CoA thiolase A, peroxisomal
				Rattus norvegicus Aldolase A, fructose-bisphosphate (Aldoa), mRNA. Length =	
1491	15511	NM_012495 NM_012498	<u>q</u> и	1442 Rattus norvegicus Aldehyde reductase 1 (low Km aldose reductase) (5.8 kb Pstl fragment, probably the functional gene) (Akr1b1), mRNA. Length = 1339	Aldolase A, fructose-bisphosphate Aldehyde reductase 1 (low Km aldose reductase) (5.8 kb Pstl fragment, probably the functional gene)

TABLE (): {	SUMMARY				Aily, Docket No. 44921-5039WG Doc. No. 1793397.1
Sequence In No.	. (Identiffer	GenBank Acc/	Model Code	Cene Name	Unigene Cluster Ville
the Manual Control of the	. iccommon	A STATE OF THE STA		Rattus norvegicus	
				Benzodiazepin receptor (peripheral) (Bzrp), mRNA.	
1494	7427	NM_012515	General	Length = 781	Benzodiazepin receptor (peripheral)
				Rattus norvegicus	
				Cholinergic receptor, muscarinic 3 (Chrm3),	
1495	24433	NM_012527	i	mRNA. Length = 3578	Cholinergic receptor, muscarinic 3
				Rattus norvegicus Creatine	
1496	4467	NM_012529	d	kinase, brain (Ckb), mRNA. Length = 1146	Creatine kinase, brain
1400	14107	14.11_012.02.0		Rattus norvegicus	
		l		Ceruloplasmin (ferroxidase)	
1497	16520	NM_012532	General	(Cp), mRNA. Length = 3700 Rattus norvegicus	Ceruloplasmin (ferroxidase)
				Angiotensin I-converting	
		1		enzyme (Dipeptidyl	
1498	225	NM 012544	x,z	carboxypeptidase 1) (Ace), mRNA. Length = 4142	Dipeptidyl carboxypeptidase 1 (Angiotensin I-converting enzyme)
1730	220	14101_012344	^,4	mistro. Congul - 4 172	variation reconverting enzyme)
				Rattus norvegicus Dopa	
				decarboxylase (aromatic L- amino acid decarboxylase)	Dopa decarboxylase (aromatic L-
1499	1431	NM 012545	General	(Ddc), mRNA. Length = 1954	
				Rattus norvegicus Early	
4500	00000	NNA 040554	I,m,v,	growth response 1 (Egr1),	Forth groudh response 4
1500	23868	NM_012551	General	mRNA. Length = 3112 Rattus norvegicus Early	Early growth response 1
			I,v,cc,	growth response 1 (Egr1),	
1500	23872	NM_012551	General	mRNA. Length = 3112	Early growth response 1
				Rattus norvegicus Early growth response 1 (Egr1),	
1500	23869	NM_012551	v,General	mRNA. Length = 3112	Early growth response 1
				Rattus norvegicus Enolase	
1501	19407	NM_012554	z	1, alpha (Eno1), mRNA. Length = 1725	Enolase 1, alpha
1501	19407	NW_012554		Rattus norvegicus Enolase	Enolase I, alpha
				1, alpha (Eno1), mRNA.	
1501	19408	NM_012554	n,s,y,z	Length = 1725 Rattus norvegicus Ets avian	Enolase 1, alpha
				erythroblastosis virus E2	
				, ,	Ets avian erythroblastosis virus E2
.=			1	progression locus 1) (Ets1),	oncogene homolog 1 (tumor
1502	21836	NM_012555	k	mRNA. Length = 4991 Rattus norvegicus Fructose-	progression locus 1)
				1,6- biphosphatase (Fbp1),	
1503	16895	NM_012558	g,s	mRNA. Length = 1357	Fructose-1,6- biphosphatase
				Rattus norvegicus Fibrinogen, gamma	
				polypeptide (Fgg), mRNA.	
1504	25317	NM_012559	bb	Length = 1358	
				Rattus norvegicus Fibrinogen, gamma	
				polypeptide (Fgg), mRNA.	
1504	6477	NM_012559	b,bb	Length = 1358	Fibrinogen, gamma polypeptide
				Rattus norvegicus	
				Fibrinogen, gamma polypeptide (Fgg), mRNA.	
1504	6478	NM_012559	bb	Length = 1358	Fibrinogen, gamma polypeptide
1505	11724	NIM 043564	L	Rattus norvegicus Follistatin	Follistatio
1505	11731	NM_012561	k	(Fst), mRNA. Length = 1035 Rattus norvegicus Group-	Follistatin
	1			specific component (vitamin	
	l			D-binding protein) (Gc),	Group-specific component (vitamin D-
1507	4254	NM_012564	а	mRNA. Length = 1676	binding protein)

TABLE 1: 8	YSSAMIMUE				* /Atty, Doctol No. 44921-5939W0 Doc. No. 1793897.1
Sequence ID No.	ldentifier:	Genbank Acc/ Ref. Seq (D	Gode Model	Gene Name	Unigane Cluster Title
1508	16026	NM_012578	r	Rattus norvegicus Histone H1-0 (H1f0), mRNA. Length = 1779	Histone H1-0
1508	16024	NM_012578	r	Rattus norvegicus Histone H1-0 (H1f0), mRNA. Length = 1779	Histone H1-0
1508	16025	NM_012578	r	Rattus norvegicus Histone H1-0 (H1f0), mRNA. Length = 1779 Rattus norvegicus Heme	Histone H1-0
1509	16080	NM_012580	g,m	oxygenase (Hmox1), mRNA. Length = 870 Rattus norvegicus Insulin-	Heme oxygenase
1510	15098	NM_012588	bb	like growth factor-binding protein (IGF-BP3) (Igfbp3), mRNA. Length = 2352	Insulin-like growth factor-binding protein (IGF-BP3)
1511	4450	NM_012592	bb	Rattus norvegicus Isovaleryl Coenzyme A dehydrogenase (Ivd), mRNA. Length = 2104	Isovaleryl Coenzyme A dehydrogenase
1511	4451	NM_012592	i,bb	Rattus norvegicus Isovaleryl Coenzyme A dehydrogenase (Ivd), mRNA. Length = 2104	Isovaleryl Coenzyme A dehydrogenase
1511	4452	NM_012592	bb	Rattus norvegicus Isovaleryl Coenzyme A dehydrogenase (Ivd), mRNA. Length = 2104	Isovaleryl Coenzyme A dehydrogenase
1512	17198	NM_012593	a,x	Rattus norvegicus Kallikrein 1, renal/pancreas/salivary (Klk1), mRNA. Length = 786	Kallikrein 1, renal/pancreas/salivary
1512	17197	NM_012593	x	Rattus norvegicus Kallikrein 1, renal/pancreas/salivary (KIk1), mRNA. Length = 786	Kallikrein 1, renal/pancreas/salivary
1513	18749	NM_012600	a,h	Rattus norvegicus Malic enzyme 1, soluble (Me1), mRNA. Length = 1761	Malic enzyme 1, soluble
				Rattus norvegicus Avian myelocytomatosis viral (v- myc) oncogene homolog	Avian myelocytomatosis viral (v-myc)
1514	2628	NM_012603	General	(Myc), mRNA. Length = 2168 Rattus norvegicus Avian	
1514	2629	NM_012603	x,General	myelocytomatosis viral (v- myc) oncogene homolog (Myc), mRNA. Length = 2168	Avian myelocytomatosis viral (v-myc) oncogene homolog
					Membrane metallo-endopeptidase
1515	16849	NM_012608	n,o,q	e) (Mme), mRNA. Length = 3243 Rattus norvegicus serine (or	(neutral endopeptidase/enkephalinase)
				cysteine) proteinase inhibitor, clade E (nexin, plasminogen activator inhibitor type 1), member 1 (Serpine1), mRNA. Length =	
1517	15540	NM_012620	General	3053 Rattus norvegicus Prolactin receptor (Prlr), mRNA.	Plasminogen activator inhibitor
1518	24568	NM_012630	General	Length = 1635 Rattus norvegicus Prolactin receptor (Prlr), mRNA.	Prolactin receptor
1518	24566	NM_012630	General	Length = 1635	Prolactin receptor

TABLE 18	SUMMARY !				Atty. Docket No. 44921-5089000 Doc. No. 1793397.
Sequence: ID No:	ldeniiier	Genbenk Acc/ Ref. Seq ID	Model Gode	Gene Name	Unigene Cluster III.le
1519	18553	NM_012631	k	Rattus norvegicus Prion protein, structural (Pmp), mRNA. Length = 765 Rattus norvegicus protein tyrosine phosphatase, non-	Prion protein, structural
1520	1844	NM_012637	General	receptor type 1 (Ptpn1), mRNA. Length = 4127	ESTs,Protein-tyrosine phosphatase
1521	24668	NM_012642	f	Rattus norvegicus Renin (Ren), mRNA. Length = 1059	Renin
1522	18632	NM_012645	а	Rattus norvegicus RT1 class lb gene (RT1Aw2), mRNA. Length = 1540	RT1 class lb gene
1523	25435	NM_012647	g	Rattus norvegicus Sodium channel, voltage-gated, type II, alpha polypeptide (Scn2a1), mRNA. Length = 8553 Rattus norvegicus	
1524	9423	NM_012649	b,cc	Ryudocan/syndecan 4 (Sdc4), mRNA. Length = 2462	Ryudocan/syndecan 4
1525	24496	NM_012654	n	Rattus norvegicus Solute carrier family 9 (sodium/hydrogen exchanger 3), antiporter 3, Na+/H+ (amiloride insensitive) (Slc9a3), mRNA. Length = 5153	Solute carrier family 9 (sodium/hydrogen exchanger 3), antiporter 3, Na+/H+ (amiloride insensitive)
1526	7101	NM_012679	x,bb, General	Rattus norvegicus Clusterin (Clu), mRNA. Length = 1638	Testostrone-repressed prostate message 2
1527	24707	NM_012693	i	Rattus norvegicus Cytochrome P450 IIA2 (Cyp2a2), mRNA. Length = 2259 Rattus norvegicus T-	Cytochrome P450 IIA2
1528	1850	NM_012696	t	kininogen, see also D11Elh1 and D11Mit8 (Kng), mRNA. Length = 1417	T-kininogen
1528	1854	NM_012696	t	Rattus norvegicus T- kininogen, see also D11Elh1 and D11Mit8 (Kng), mRNA. Length = 1417	K-kininogen, differential splicing leads to HMW Kngk,T-kininogen
1529	1603	NM_012697	General	Rattus norvegicus Organic cation transporter (Slc22a1), mRNA. Length = 1882 Rattus norvegicus	Organic cation transporter
1530	1372	NM_012734	u	Hexokinase 1 (Hk1), mRNA. Length = 3653 Rattus norvegicus Pyruvate	Hexokinase 1
1531	1478	NM_012744	bb, General	carboxylase (Pc), mRNA. Length = 3945 Rattus norvegicus Signal	Pyruvate carboxylase
1532	343	NM_012747	h,t	transducer and activator of transcription 3 (Stat3), mRNA. Length = 2924	Signal transducer and activator of transcription 3
1533	8829	NM_012749	General	Rattus norvegicus Nucleolin (Ncl), mRNA. Length = 2142	Nucleolin
1534	20828	NM_012752	General	Rattus norvegicus CD24 antigen (Cd24), mRNA. Length = 1703 Rattus norvegicus CD24	CD24 antigen
1534	20829	NM_012752	i,General	antigen (Cd24), mRNA. Length = 1703	CD24 antigen

TABLE 1: 8	UMINARY	i i i i i i i i i i i i i i i i i i i			Atty, Docket No. 44921-5039770 Doc. No. 1793397.1
Sequence	Octors (MT) con	ConBank Acc Ref. Seq ID	Model Gode	Cono Namo	Unicano Civeter Titlo
1534	20830	NM_012752	i,General	Rattus norvegicus CD24 antigen (Cd24), mRNA. Length = 1703	CD24 antigen
:				Rattus norvegicus Insulin- like growth factor 2 receptor (Igf2r), mRNA. Length =	
1535	15174	NM_012756	b	8810 Rattus norvegicus Lost on	Insulin-like growth factor 2 receptor
1536	21685	NM_012760	j,m,n	transformation 1 (Lot1), mRNA. Length = 5028	Lost on transformation 1
1537	18068	NM 012762	t	Rattus norvegicus Interleukin 1beta converting enzyme (Casp1), mRNA. Length = 1209	Interleukin 1beta converting enzyme
1538	1246	NM 012770	a,General		Guanylate cyclase, soluble, beta 2 (GTP pyrophosphate - lyase)
1539	1348	NM_012776	f	Rattus norvegicus adrenergic receptor kinase, beta 1 (Adrbk1), mRNA. Length = 2683	G-protein-linked receptor kinase (beta adrenergic receptor kinase 1)
1540	18135	NM 012791	w	Rattus norvegicus dual- specificity tyrosine-(Y)- phosphorylation regulated kinase 1a (Dyrk1a), mRNA. Length = 2840	Dual Specificity Yak1-related kinase,ESTs
1541	16947	NM_012793	p,bb	Rattus norvegicus Guanidinoacetate methyltransferase (Gamt), mRNA. Length = 924	Guanidinoacetate methyltransferase
				Rattus norvegicus glutathione S-transferase, theta 2 (Gstt2), mRNA.	alidathian Changain that 2
1542	960	NM_012796	u	Length = 1258 Rattus norvegicus MAL protein gene (Mal), mRNA.	glutathione S-transferase, theta 2
1543	260	NM_012798	f,u ·	Length = 2268 Rattus norvegicus Protein C	MAL protein gene
1544	556	NM_012803	d	(Proc), mRNA. Length = 1543	Protein C
1545	21729	NM_012804	q	Rattus norvegicus ATP- binding cassette, sub-family D (ALD), member 3 (Abcd3), mRNA. Length = 3324	ATP-binding cassette, sub-family D (ALD), member 3
1343	21125	14141 0 12004	4	Rattus norvegicus alpha- methylacyl-CoA racemase (Amacr), mRNA. Length =	(ACD), Member 0
1546	15032	NM_012816	General	1504 Rattus norvegicus Insulin- like growth factor-binding protein 5 (Igfbp5), mRNA.	Methylacyl-CoA racemase alpha Insulin-like growth factor-binding
1547	24895	NM_012817	General	Length = 1630 Rattus norvegicus Annexin A3 (Anx3), mRNA. Length =	protein 5 ESTs, Weakly similar to LURT3
1548	18109	NM_012823	u,General	1454 Rattus norvegicus ATP- binding cassette, sub-family C (CFTR/MRP), member 2	annexin III - rat [R.norvegicus]
1549	373	NM_012833	h,l,q, General	(Abcc2), mRNA. Length = 4918 Rattus norvegicus Cystatin	Canalicular multispecific organic anion transporter
1550	2855	NM_012838	е	beta (Cstb), mRNA. Length = 590	Cystatin beta

TRABLE 1:	SUMMARY				Air. Docket No. 44921-5039WG Doc. No. 1793897.
Sequence ID No.	Midentifier	Genbenk Ace Ref. Seq.(D)	Mode) Gode	Gene Name	Uilgene Cluster Title
				Rattus norvegicus Cytochrome C, expressed in somatic tissues (Cycs),	Cytochrome C, expressed in somatic
1551	11136	NM_012839	s	mRNA. Length = 318 Rattus norvegicus Epidermal	tissues
1552	20885	NM_012842	а	growth factor (Egf), mRNA. Length = 4801 Rattus norvegicus Epidermal	Epidermal growth factor
1552	20884	NM_012842	a,bb	growth factor (Egf), mRNA. Length = 4801 Rattus norvegicus	Epidermal growth factor
1553	18770	NM 012857	e	Lysosomal associated membrane protein 1 (120 kDa) (Lamp1), mRNA. Length = 2006	Lysosomal associated membrane protein 1 (120 kDa)
1550	10770	1411_012331		Rattus norvegicus O6- methylguanine-DNA methyltranferase (Mgmt).	ESTs, Weakly similar to S21348 probable pol polyprotein-related protein 4 - rat [R.norvegicus], O6-
1554	20674	NM_012861	i	mRNA. Length = 812 Rattus norvegicus Matrix Gla	methylguanine-DNA methyltranferase
1555	13151	NM_012862	a,r, General	protein (Mgp), mRNA. Length = 521	Matrix Gla protein
				Rattus norvegicus tumor necrosis factor receptor superfamily, member 11b (osteoprotegerin) (Tnfrsf11b), mRNA. Length =	
1556	24617	NM_012870	General	2432 Ribosomal protein L39	Osteoprotegerin
1557	20945	NM 012875	a,v	(Rpl39), mRNA. Length = 324	Ribosomal protein L39
1558	15872	NM 012879	o,r	Rattus norvegicus Solute carrier family 2 A2 (gkucose transporter, type 2) (Slc2a2), mRNA. Length = 2573	Solute carrier family 2 A2 (gkucose transporter, type 2)
				Rattus norvegicus Superoxide dismutase 3 (Sod3), mRNA. Length =	
1559	495	NM_012880	z	1729 Rattus norvegicus	Superoxide dimutase 3
1559	494	NM_012880	c_	Superoxide dismutase 3 (Sod3), mRNA. Length = 1729	Superoxide dimutase 3
			d,u,	Rattus norvegicus Sialoprotein (osteopontin) (Spp1), mRNA. Length =	
1560	23651	NM_012881	General	1457 Rattus norvegicus Acyl-Coa	Sialoprotein (osteopontin) EST, Moderately similar to ACDV RATACYL-COA DEHYDROGENASE,
1562	19477	NM_012891	q	dehydrogenase, Very long chain (Acadvl), mRNA. Length = 2117	VERY-LONG-CHAIN SPECIFIC, MITOCHONDRIAL PRECURSOR [R.norvegicus]
				Rattus norvegicus aminolevulinate,delta- ,dehydratase (Alad), mRNA.	Delta - aminolevulinic acid
1563	18564	NM_012899		Length = 1116 Rattus norvegicus Annexin 1	dehydratase
1564	7197	NM_012904	f,r,cc,Gen eral	(p35) (Lipocortin 1) (Anxa1), mRNA. Length = 1402	Annexin 1 (p35) (Lipocortin 1)
1564	7196	NM_012904	v,cc, General	Rattus norvegicus Annexin 1 (p35) (Lipocortin 1) (Anxa1), mRNA. Length = 1402	Annexin 1 (p35) (Lipocortin 1)
1565	20202	NM 012909	b,r	Rattus norvegicus Aquaporin 2 (Aqp2), mRNA. Length = 939	Aquaporin 2
					

TABLE 1:	SUMMARY				Attyr. Docket No. 44921-5039W0 Dock No. 1793397.1
Government.		GenBank Acc/	Madal		
Economica ID No.	ldentifier	Ref. Seq ID		Cene Name	Unigene Gluster Tille
				Rattus norvegicus Arrestin, beta 2 (Arrb2), mRNA.	
1566	16581	NM_012911	c,j	Length = 1758 Rattus norvegicus Arrestin,	Arrestin, beta 2
1566	16582	NM_012911	С	beta 2 (Arrb2), mRNA. Length = 1758	Arrestin, beta 2
1567	24431	NM 012912	General	Rattus norvegicus Activating transcription factor 3 (Atf3), mRNA. Length = 1893	Activating transcription factor 3
4500	10110	NIM 049042		Rattus norvegicus ATPase, Na+K+ transporting, beta polypeptide 3 (Atp1b3),	ATPase, Na+K+ transporting, beta
1568	18118	NM_012913	р	mRNA. Length = 1818 Rattus norvegicus ATPase inhibitor (rat mitochondrial	polypeptide 3
1569	6108	NM_012915	n	IF1 protein) (Atpi), mRNA. Length = 833	ATPase inhibitor (rat mitochondrial IF1 protein)
1570	20757	NM 012923	c,i,aa	Rattus norvegicus Cyclin G1 (Ccng1), mRNA. Length = 3169	Cyclin G1
				Rattus norvegicus Cyclin G1 (Ccng1), mRNA. Length =	
1570	20755	NM_012923	i	3169 Rattus norvegicus CD59	Cyclin G1
4574	0000	NIN 04000E		antigen (Cd59), mRNA.	CD50 antigon
1571	2830	NM_012925	<u> </u>	Length = 1523 Rattus norvegicus CD59	CD59 antigen
1571	2831	NM_012925	f	antigen (Cd59), mRNA. Length = 1523	CD59 antigen
1572	1977	NM_012930	q	Rattus norvegicus Carnitine palmitoyltransferase 2 (Cpt2), mRNA. Length = 2296	Carnitine palmitoyltransferase 2
				Rattus norvegicus v-crk- associated tyrosine kinase substrate (Crkas), mRNA.	v-crk-associated tyrosine kinase
1573	18694	NM_012931	j,l,m,z	Length = 3335 Rattus norvegicus Crystallin,	substrate
1574	13723	NM_012935	n	alpha polypeptide 2 (Cryab), mRNA. Length = 528	Crystallin, alpha polypeptide 2,ESTs
				Rattus norvegicus Cathepsin H (Ctsh), mRNA. Length =	
1575	9109	NM_012939	j,y,z	1362 Rattus norvegicus Cathepsin	Cathepsin H
1575	19398	NM_012939	aa	H (Ctsh), mRNA. Length = 1362	 EST
				Rattus norvegicus Diphtheria toxin receptor(heparin binding epidermal growth	Diphtheria toxin receptor (heparin
1576	223	NM_012945	b,cc	factor - like growth factor) (Dtr), mRNA. Length = 1550	binding epidermal growth factor - like growth factor)
1577	15058	NM_012950	cc	Rattus norvegicus Thrombin receptor (F2r), mRNA. Length = 3418	Thrombin receptor
1579	19111	NM 012963	g	Rattus norvegicus High mobility group 1 (Hmg1), mRNA, Length = 1225	High mobility group 1
	70171	1	3	Rattus norvegicus Hyaluronan mediated motility	
1580	19374	NM_012964	x	receptor (RHAMM) (Hmmr), mRNA. Length = 2049 Rattus norvegicus	Hyaluronan mediated motility receptor (RHAMM)
4504	0554			Intercellular adhesion molecule 1 (Icam1), mRNA.	
1581	2554	NM_012967	Įt.	Length = 2602	Intercellular adhesion molecule 1

TABLE 1: S	UMMARY		J.		Atty. Docto: No. 44221-5039W0 Doc. No. 1793397.1
Sequence ID No.	ldentifer	GenBenk Acc/ Ref. Seq ID	Model Code	Gene Name	Unigene Guster Title
1581	2555	NM_012967	t,cc, General	Rattus norvegicus Intercellular adhesion molecule 1 (Icam1), mRNA. Length = 2602 Rattus norvegicus Potassium	Intercellular adhesion molecule 1
1582	24528	NM_012973	c	(K+) channel protein, slowly activating (lsk) (Kcne1), mRNA. Length = 585 Rattus norvegicus Lectin,	Potassium (K+) channel protein, slowly activating (Isk)
1583	956	NM_012976	С	galactose binding, soluble 5 (Galectin-5) (Lgals5), mRNA. Length = 872	Lectin, galactose binding, soluble 9 (Galectin-9)
1584	16417	NM_012991	g	Rattus norvegicus Nucleoprotein 50kD (Nup50), mRNA. Length = 3027	Nuclear pore associated protein
1585	17393	NM_012992	d	Rattus norvegicus Nucleoplasmin-related protein (Nuclear protein B23 (Npm1), mRNA. Length = 1232	Nucleoplasmin-related protein (Nuclear protein B23
1586	23544	NM_013013	s	Rattus norvegicus Prosaposin (sulfated glycoprotein, sphingolipid hydrolase activator) (Psap), mRNA. Length = 2175	Prosaposin (sulfated glycoprotein, sphingolipid hydrolase activator)
1587	1588	NM 013026	k	Rattus norvegicus Syndecan 1 (Sdc1), mRNA. Length = 2410	Syndecan 1
1588	17894	NM 013027	m ·	Rattus norvegicus Selenoprotein W muscle 1 (Sepw1), mRNA. Length = 664	Selenoprotein W muscle 1
1589	18300	NM 013030	s,v, General	Rattus norvegicus Solute carrier family 17 (sodium/hydrogen exchanger), member 2 (Slc34a1), mRNA. Length = 2440	Rattus norvegicus mRNA for NaPi-2
				Rattus norvegicus Solute carrier family 17 (sodium/hydrogen exchanger), member 2 (Sic34a1), mRNA. Length =	Solute carrier family 17 (sodium/hydrogen exchanger),
1589	18076	NM_013030	g,s,z	2440 Rattus norvegicus Solute carrier family 17 (sodium/hydrogen exchanger), member 2 (Slc34a1), mRNA. Length =	member 2 Rattus norvegicus mRNA for NaPi-2 alpha, complete cds,Solute carrier family 17 (sodium/hydrogen
1589	18078	NM_013030	S	2440 Rattus norvegicus Solute carrier family 17 (sodium/hydrogen exchanger), member 2	exchanger), member 2 Solute carrier family 17
1589	18077	NM_013030	e,s,z	(Slc34a1), mRNA. Length = 2440 Rattus norvegicus ATP-	(sodium/hydrogen exchanger), member 2
1591	730	NM_013040	w	binding cassette, sub-family C (CFTR/MRP), member 9 (Abcc9), mRNA. Length = 5000	Sulfonylurea receptor 2

TABLE 1: S	UMIMARY				Atty, Docket No. 44921-5039W0 Doc. No. 1793397.1
Sequence ID No.	l Clean Miles	©елВалк Acc/ Ref. Seq ID	Model Code	Gene Name	Unigene Cluster tittle
1592	17401	NM_013043	i,o, General	Rattus norvegicus Transforming growth factor beta stimulated clone 22 (Tgfb1i4), mRNA. Length = 1666	Transforming growth factor beta stimulated clone 22
1593	16684	NM_013052	General	Rattus norvegicus Tyrosine 3 monooxygenase/tryptophan 5-monooxygenase activation protein, eta polypeptide (Ywhah), mRNA. Length = 1689	
1594	14421	NM_013053	u	Rattus norvegicus Tyrosine 3 monooxygenase/tryptophan 5-monooxygenase activation protein, theta polypeptide (Ywhaq), mRNA. Length = 2099	
1595	15254	NM_013058	k	Rattus norvegicus Inhibitor of DNA binding 3, dominant negative helix-loop-helix protein (Id3), mRNA. Length = 568	Inhibitor of DNA binding 3, dominant negative helix-loop-helix protein
1596	14997	NM_013059	s,z	Rattus norvegicus Tissue- nonspecific ALP alkaline phosphatase (AlpI), mRNA. Length = 2415 Rattus norvegicus Tissue-	Tissue-nonspecific ALP alkaline phosphatase
1596	14996	NM_013059	General	nonspecific ALP alkaline phosphatase (AlpI), mRNA. Length = 2415	Tissue-nonspecific ALP alkaline phosphatase
4507	25676	NM 013069	aa	Rattus norvegicus CD74 antigen (invariant polpypeptide of major histocompatibility class II antigen-associated) (Cd74), mRNA. Length = 1150	
1597	16924	NM 013069	0	Rattus norvegicus CD74 antigen (invariant polpypeptide of major histocompatibility class II antigen-associated) (Cd74), mRNA. Length = 1150	CD74 antigen (invariant polpypeptide of major histocompatibility class II antigen-associated)
1598	24748	NM 013070	h,q	Rattus norvegicus Utrophin (Utrn), mRNA. Length = 10,705	Utrophin
1599	1529	NM_013082		Rattus norvegicus Ryudocan/syndecan 2 (Sdc2), mRNA. Length = 2153	Ryudocan/syndecan 2
1600	1521	NM 013091	j,l,z, General	Rattus norvegicus Tumor necrosis factor receptor superfamily, member 1a (Tnfr1), mRNA. Length = 2130	Tumor necrosis factor receptor
1601	1685	NM_013096	c,aa	Rattus norvegicus Hemoglobin, alpha 1 (Hba1), mRNA. Length = 556	Hemoglobin, alpha 1
1601	26150	NM_013096	c,i	Rattus norvegicus Hemoglobin, alpha 1 (Hba1), mRNA. Length = 556 Rattus norvegicus	
1601	1688	NM_013096	p	Hemoglobin, alpha 1 (Hba1), mRNA. Length = 556	Hemoglobin, alpha 1

Table 1: 8	UMMARY				#Affy, Docket No. 44921-5039W0 Doc. No. 1793897.1
Sequence ID No.	licalifier .		Model Code	Cene Name	Unigane Civeter Tille
1601	1689	NM_013096	с,р	Rattus norvegicus Hemoglobin, alpha 1 (Hba1), mRNA. Length = 556	Hemoglobin, alpha 1
1601	1684	NM_013096	c,s,aa	Rattus norvegicus Hemoglobin, alpha 1 (Hba1), mRNA. Length = 556	Hemoglobin, alpha 1
1602	20886	NM 013097	u,x,bb	Rattus norvegicus Deoxyribonuclease I (Dnase1), mRNA. Length = 1143	Deoxyribonuclease I
1602	20887	NM_013097	u,x,bb	Rattus norvegicus Deoxyribonuclease I (Dnase1), mRNA. Length = 1143	Deoxyribonuclease I
1603	1321	NM_013098	с	Rattus norvegicus Glucose-6- phosphatase (G6pc), mRNA. Length = 2237	Glucose-6-phosphatase
1604	15296	NM 013102	I,m	Rattus norvegicus FK506- binding protein 1 (12kD) (Fkbp1a), mRNA. Length = 554	FK506-binding protein 1 (12kD)
1004				Rattus norvegicus ATPase Na+/K+ transporting beta 1 polypeptide (Atp1b1),	ATPase Na+/K+ transporting beta 1
1606	23709	NM_013113	o,s,z,aa	mRNA. Length = 2528 Rattus norvegicus ATPase Na+/K+ transporting beta 1	polypeptide
1606	23711	NM_013113	p	polypeptide (Atp1b1), mRNA. Length = 2528 Rattus norvegicus ATPase	ATPase Na+/K+ transporting beta 1 polypeptide
1606	23710	NM_013113	s	Na+/K+ transporting beta 1 polypeptide (Atp1b1), mRNA. Length = 2528	ATPase Na+/K+ transporting beta 1 polypeptide
1607	1976	NM_013118	u	Rattus norvegicus Guanylate cyclase activator 2 (guanylin) (Guca2a), mRNA. Length = 567	Guanylate cyclase activator 2 (guanylin)
1609	870	NM_013130	h	Rattus norvegicus MAD (mothers against decapentaplegic, Drosophila) homolog 1 (Madh1), mRNA. Length = 2002	MAD (mothers against decapentaplegic, Drosophila) homolog
1610	16650	NM_013132	u,General	Rattus norvegicus Annexin V (Anx5), mRNA. Length = 1417	Annexin V
1611	650	NM_013134	h	Rattus norvegicus 3-hydroxy- 3-methylglutaryl-Coenzyme A reductase (Hmgcr), mRNA. Length = 2664	3-hydroxy-3-methylglutaryl-Coenzyme A reductase
1611	651	NM_013134	h,j,l	Rattus norvegicus 3-hydroxy- 3-methylglutaryl-Coenzyme A reductase (Hmgcr), mRNA. Length = 2664	3-hydroxy-3-methylglutaryl-Coenzyme A reductase
4040	4740	NIM 042422	Concret	Rattus norvegicus Inositol 1, 4, 5-triphosphate receptor 3 (Itpr3), mRNA. Length =	Inocital 1 A 5 triphocoboto recenter 2
1612	1712	NM_013138	General o,v,	8806 Rattus norvegicus Insulin- like growth factor binding protein 1 (Igfbp1), mRNA.	Inositol 1, 4, 5-triphosphate receptor 3 Insulin-like growth factor binding
1613	16982	NM_013144	General	Length = 1500	protein 1

TABLE 1: S	WMMARY				ZXIY. Docket No. 44921-5039W0 Doc. No. 1793397.1
Sequence ID No.	ldentifier	Genbank Acc Ref. Seq ID	Model Gode	Cene Name	Unitgene Cluster Title
				Rattus norvegicus CCAAT/enhancerbinding, protein (C/EBP) delta	
1614	21683	NM_013154	t,cc, General	(Cebpd), mRNA. Length =	CCAAT/enhancerbinding, protein (C/EBP) delta
				Rattus norvegicus CCAAT/enhancerbinding, protein (C/EBP) delta (Cebpd), mRNA. Length =	CCAAT/enhancerbinding, protein
1614	21682	NM_013154	сс	1200 Rattus norvegicus Cathepsin	(C/EBP) delta
1615	3431	NM_013156	b,g,n	L (CtsI), mRNA. Length = 1386 Rattus norvegicus Cathepsin	Cathepsin L
1615	25567	NM_013156	v,General	L (Ctsl), mRNA. Length = 1386 Rattus norvegicus Cathepsin	
1015	0.400	NINA 040450	C = = = = = 1	L (Ctsl), mRNA. Length =	Cathonain
1615	3430	NM_013156	General	Rattus norvegicus Insulin	Cathepsin L
1616	1309	NM_013159	l _w	degrading enzyme (Ide), mRNA. Length = 4276	Insulin degrading enzyme
1010	1303	14M_010103		Rattus norvegicus Insulin	modified and grading only mo
1616	1310	NM 013159	l _w	degrading enzyme (Ide), mRNA. Length = 4276	Insulin degrading enzyme
				Rattus norvegicus Transforming growth factor, beta 3 (Tgfb3), mRNA.	
1617	21723	NM_013174	w_	Length = 2633 Rattus norvegicus Protein	Transforming growth factor, beta 3
1618	1314	NM_013181	m	kinase, cAMP dependent, regulatory, type 1 (Prkar1a), mRNA. Length = 1433	Protein kinase, cAMP dependent, regulatory, type 1
1619	17357	NM 013183	p,bb, General	Rattus norvegicus Meprin 1 beta (Mep1b), mRNA. Length = 2290	Meprin 1 beta
				Rattus norvegicus Phosphofructokinase, liver, B-type (Pfkl), mRNA. Length	·
1620	1300	NM_013190	у	= 2740 Rattus norvegicus	Phosphofructokinase, liver, B-type
1621	16448	NM_013197	С	Aminolevulinate synthase 2, delta (Alas2), mRNA. Length = 1899	Aminolevulinate synthase 2, delta
				Rattus norvegicus Carnitine palmitoyltransferase 1 beta, muscle isoform (Cpt1b),	Carnitine palmitoyltransferase 1 beta,
1622	20856	NM_013200	b	mRNA. Length = 2826	muscle isoform Rattus norvegicus brain cytosolic acyl
1623	397	NM_013214	f	Rattus norvegicus acyl-CoA hydrolase (RBACH), mRNA. Length = 1523	coenzyme A thioester hydrolase mRNA, complete cds,acyl-CoA hydrolase
				Rattus norvegicus aflatoxin B1 aldehyde reductase (AFAR), mRNA, Length =	
1624	20864	NM_013215	g,n,y	1272	aflatoxin B1 aldehyde reductase
1625	20728	NM_013217	v	Rattus norvegicus afadin (AF 6), mRNA. Length = 5957 Rattus norvegicus	afadin
1626	1396	NM 013222	l _i	augmenter of liver regeneration (ALR), mRNA. Length = 1226	augmenter of liver regeneration
1020	1000	INIT O I DEZE	'	Rattus norvegicus ribosomal protein S26 (Rps26), mRNA.	assumed of liver regeneration
1627	815	NM_013224	w	Length = 435	ribosomal protein S26

TABLE 18	SUMMARY		- 34.4		/ // // // Dockei No. 44921-5039WC Doc. No. 1793897-1
Sequenc (D No.	i. IdenMiler	GenBank Acc/ Ref. Seq ID	Model Code	Gene Name	Unigene Civister Title
				Rattus norvegicus ribosomal protein L32 (Rpl32), mRNA.	
1628	18305	NM_013226	v	Length = 465 Rattus norvegicus Acyl-	
1629	21078	NM_016986	d	Coenzyme A dehydrogenase, C-4 to C-12 straight-chain (Acadm), mRNA. Length = 1866	Acyl-Coenzyme A dehydrogenase, C- 4 to C-12 straight-chain
				Rattus norvegicus Acid phosphatase 2, lysozymal (Acp2), mRNA. Length =	
1630	24649	NM_016988	v	2009 Rattus norvegicus adenylate	Acid phosphatase 2, lysozymal
1631	15239	NM_016989	q,w	cyclase activating polypeptide 1 (Adcyap1), mRNA. Length = 2681	R.norvegicus (Sprague Dawley) ribosomal protein L15 mRNA
4622	45	NM 016996	General	Rattus norvegicus Calcium- sensing receptor (hypocalciuric hypercalcemia 1, severe neonatal hyperparathyroidism) (Casr), mRNA. Length = 4113	Calcium-sensing receptor (hypocalciuric hypercalcemia 1, severe neonatal hyperparathyroidism)
1632	45	NM_0 16996	General		severe neonatal hyperparatity outsing
1633	20714	NM_016999	t	Rattus norvegicus Cytochrome P450, subfamily IVB, polypeptide 1 (Cyp4b1), mRNA. Length = 2462	
1633	20713	NM_016999	t	Rattus norvegicus Cytochrome P450, subfamily IVB, polypeptide 1 (Cyp4b1), mRNA. Length = 2462	Cytochrome P450, subfamily IVB, polypeptide 1
1633	20711	NM 016999	g,t	Rattus norvegicus Cytochrome P450, subfamily IVB, polypeptide 1 (Cyp4b1), mRNA. Length = 2462	Cytochrome P450, subfamily IVB, polypeptide 1
1633	20715	NM_016999	q,t	Rattus norvegicus Cytochrome P450, subfamily IVB, polypeptide 1 (Cyp4b1), mRNA. Length = 2462	
			e,n,p,	Rattus norvegicus Diaphorase (NADH/NADPH) (Dia4), mRNA. Length =	
1634	1698	NM_017000	General	1396	Diaphorase (NADH/NADPH)
1635	1399	NM_017006	h,n, General	Rattus norvegicus Glucose-6 phosphate dehydrogenase (G6pd), mRNA. Length = 2324	Glucose-6-phosphate dehydrogenase
1637	18989	NM_017013	n	Rattus norvegicus Glutathione-S-transferase, alpha type (Yc?) (Gsta2), mRNA. Length = 830	Glutathione-S-transferase, alpha type (Yc?)
1638	21013	NM_017014	e,f	Rattus norvegicus Glutathione-S-transferase, mu type 2 (Yb2) (Gstm2), mRNA. Length = 1055	Glutathione-S-transferase, mu type 2 (Yb2)
1638	21015	NM_017014	e,General	Rattus norvegicus Glutathione-S-transferase, mu type 2 (Yb2) (Gstm2), mRNA. Length = 1055	Glutathione-S-transferase, mu type 2 (Yb2)

TABLE 1: S	UMMARY				AMy. Docket No. 44921-5039W0 Dock No. 1793897.1
Sequence ID No:	loealijars	Conbant Acc Ref Seq ID	Model Code	Cene Name	ee ke 11999 in Uilijane Glyslat Tille
	licentifies.		<u> </u>	Rattus norvegicus Potassium inwardly-rectifying channel, subfamily J (Kcnj1), mRNA.	Potassium inwardly-rectifying channel,
1639	11836	NM_017023	b	Length = 2069 Rattus norvegicus Potassium	subfamily J
1639	5475	NM 017023	b	inwardly-rectifying channel, subfamily J (Kcnj1), mRNA. Length = 2069	ESTs,Potassium inwardly-rectifying channel, subfamily J
				Rattus norvegicus Potassium inwardly-rectifying channel, subfamily J (Kcnj1), mRNA.	
1639	25546	NM_017023	b,bb	Length = 2069 Rattus norvegicus Lactate	
1640	17807	NM_017025	i,General	dehydrogenase A (Ldha), mRNA. Length = 1609 Rattus norvegicus Protein	Lactate dehydrogenase A
1641	24597	NM_017040	u	phosphatase 2 (formerly 2A), catalytic subunit, beta isoform (Ppp2cb), mRNA. Length = 1843	Protein phosphatase 2 (formerly 2A), catalytic subunit, beta isoform
				Rattus norvegicus Solute carrier family 4, member 2, anion exchange protein 2 (Slc4a2), mRNA. Length =	Solute carrier family 4, member 2,
1642	24696	NM_017048	f,j,z	4057 Rattus norvegicus Solute	anion exchange protein 2
1643	24695	NM_017049	u	carrier family 4, member 3, anion exchange protein 3 (Slc4a3), mRNA. Length = 3877	Solute carrier family 4, member 3, anion exchange protein 3
				Rattus norvegicus Superoxide dismutase 1, soluble (Sod1), mRNA.	
1644	20876	NM_017050	j,n,z	Length = 650 Rattus norvegicus Bcl2-	Superoxide dimutase 1, soluble
1645	910	NM_017059	f,I,m	associated X protein (Bax), mRNA. Length = 579	Bcl2-associated X protein
				Rattus norvegicus Bcl2- associated X protein (Bax),	D-10isted V seetsin
1645	912	NM_017059	<u> </u>	mRNA. Length = 579 Rattus norvegicus Lysyl oxidase (Lox), mRNA.	Bcl2-associated X protein
1646	1946	NM_017061	h	Length = 4557 Rattus norvegicus Lysyl	Lysyl oxidase
1646	1942	NM_017061	t,General	oxidase (Lox), mRNA. Length = 4557 Rattus norvegicus Lysyl	Lysyl oxidase
1646	1943	NM_017061	t	oxidase (Lox), mRNA. Length = 4557	Lysyl oxidase
				Rattus norvegicus Pleiotrophin (Heparine binding factor, Hbnf, in the	
1647	6062	NM_017066	d	= 1246 Rattus norvegicus	Pleiotrophin (Heparine binding factor, Hbnf, in the mouse)
				Lysosomal-associated membrane protein 2 (Lamp2), mRNA. Length =	Lysosomal-associated membrane
1648	6654	NM_017068	w	Rattus norvegicus Glutamine synthetase (glutamate-	protein 2
1649	11153	NM_017073	s	ammonia ligase) (Glul), mRNA. Length = 2793	Glutamine synthetase (glutamate- ammonia ligase)

TABLE 1: 8	MIMARY				Atty, Docket No. 44921-5039W0 Doc. No. 1793897.1
Sequence ID No.	Oslara Wilesz	GenBenk Acc Ref Sco ID	Model Gode	Gene Name	
	i identitie			Rattus norvegicus Tumor- associated glycoprotein pE4 (Tage4), mRNA. Length =	Commission — A management to produce the state of the sta
1650	923	NM_017076	General	2171 Rattus norvegicus CD1D	Tumor-associated glycoprotein pE4
1651	1523	NM_017079	s	antigen (Cd1d), mRNA. Length = 1835 Rattus norvegicus	CD1D antigen
1652	23660	NM 017080	s	Hydroxysteroid dehydrogenase, 11 beta type 1 (Hsd11b1), mRNA. Length = 1265	Hydroxysteroid dehydrogenase, 11 beta type 1
1002	2000	0	b,d,	Rattus norvegicus Hydroxysteroid dehydrogenase, 11 beta type 2 (Hsd11b2), mRNA. Length	Hydroxysteroid dehydrogenase, 11
1653	275	NM_017081	General	= 1864 Rattus norvegicus Urmodulin	beta type 2
1654	16211	NM_017082	j,s,z	(Tamm-Horsfall protein) (Umod), mRNA. Length = 2227	Urmodulin (Tamm-Horsfall protein)
1655	1552	NM_017084	j	Rattus norvegicus Glycine methyltransferase (Gnmt), mRNA. Length = 988	Glycine methyltransferase
1655	1550	NM_017084	у	Rattus norvegicus Glycine methyltransferase (Gnmt), mRNA. Length = 988	Glycine methyltransferase
1656	22552	NM 017087	a,k,x	Rattus norvegicus Small proteoglycan I (biglycan), bone (BSPG1) (bone/cartilage proteclycan 1 precursor) (Bgn), mRNA. Length = 2446	Small proteoglycan I (biglycan), bone (BSPG1) (bone/cartilage proteclycan 1 precursor)
				Rattus norvegicus guanylate cyclase 1, soluble, alpha 3 (Gucy1a3), mRNA. Length = 4775	Guanylate cyclase, soluble, alpha 1
1657	8888	NM_017090	m	Rattus norvegicus Growth hormone receptor (Ghr),	(GTP pyrophosphate - lyase)
1658	10887	NM_017094	a,General	mRNA. Length = 2950 Rattus norvegicus	Growth hormone receptor
1659	4393	NM_017101	а,у	Length = 743	Peptidylprolyl isomerase A (cyclophilin A)
				Rattus norvegicus solute carrier family (organic anion transporter) member 1 (Slc21a1), mRNA. Length =	solute carrier family (organic anion
1660	24770	NM_017111	đ	2758 Rattus norvegicus granulin	transporter) member 1
1661	20745	NM_017113	e	(Grn), mRNA. Length = 2113 Rattus norvegicus granulin	
1661	20746	NM_017113	а	(Grn), mRNA. Length = 2113 Rattus norvegicus	granulin
1662	1375	NM_017122	w	hippocalcin (Hpca), mRNA. Length = 1561 Rattus norvegicus CD37	hippocalcin
1663	12903	NM_017124	k	antigen (Cd37), mRNA. Length = 1158 Rattus norvegicus laminin	CD37 antigen
1664	24885	NM_017138	r	receptor 1 (Lamr1), mRNA. Length = 1018	laminin receptor 1
1664	24886	NM_017138	d,q	Rattus norvegicus laminin receptor 1 (Lamr1), mRNA. Length = 1018	laminin receptor 1

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TABLE 1: 8					Any, Docket No. 44921-5039Wo Doc. No. 1793897.1
Sequence ID No.	ldemiller	Genbenk Acc/ Ref. Seq ID	Model Code	Cene Name	Unigano Clustor Tillo
1665	15363	NM_017147	n,u	Rattus norvegicus cofilin 1, non-muscle (Cf11), mRNA. Length = 1039	cofilin 1, non-muscle
1666	13392	NM_017148	u,General	Rattus norvegicus cysteine rich protein 1 (Csrp1), mRNA. Length = 1403 Rattus norvegicus ribosomal	cysteine rich protein
1667	5351	NM_017150	q	protein L29 (Rpl29), mRNA. Length = 630 Rattus norvegicus ribosomal	ribosomal protein L29
1668	16954	NM_017151	a,n	protein S15 (Rps15), mRNA. Length = 487 Rattus norvegicus ribosomal	ribosomal protein S15
1669	21643	NM_017152	g	protein S17 (Rps17), mRNA. Length = 466 Rattus norvegicus ribosomal	ribosomal protein S17
1670	1694	NM_017153	a,q	protein S3a (Rps3a), mRNA. Length = 880 Rattus norvegicus ribosomal	ribosomal protein S3a
1671	17104	NM_017160	bb, General	protein S6 (Rps6), mRNA. Length = 801 Rattus norvegicus ribosomal	ribosomal protein S6
1671	17106	NM_017160	u	protein S6 (Rps6), mRNA. Length = 801 Rattus norvegicus ribosomal	ribosomal protein S6
1671	17107	NM_017160	d,e	protein S6 (Rps6), mRNA. Length = 801	ribosomal protein S6
1672	17686	NM_017165	n,q	Rattus norvegicus glutathione peroxidase 4 (Gpx4), mRNA. Length = 872	
				Rattus norvegicus Leukemia- associatedcytosolic phosphoprotein stathmin (Lap18), mRNA. Length =	Leukemia-associated cytosolic
1673	20702	NM_017166	С	1054 Rattus norvegicus choline/ethanolamine kinase	phosphoprotein stathmin
1674	3513	NM_017177	r	(Chetk), mRNA. Length = 1679 Rattus norvegicus T-cell	choline/ethanolamine kinase
1675	19031	NM_017180	v,General	death associated gene (Tdag), mRNA. Length =	T-cell death associated gene
				Rattus norvegicus high mobility group box 2 (Hmgb2), mRNA. Length =	
1676	15437	NM_017187	x,z	Rattus norvegicus high mobility group box 2	high mobility group protein 2
1676	15433	NM_017187	у	(Hmgb2), mRNA. Length = 1072 Rattus norvegicus high	high mobility group protein 2
1676	15434	NM_017187	x,z	mobility group box 2 (Hmgb2), mRNA. Length = 1072	high mobility group protein 2
				Rattus norvegicus Myelin- associated glycoprotein (Mag), mRNA. Length =	
1677	24437	NM_017190	p	Rattus norvegicus kynurenine aminotransferase	Myelin-associated glycoprotein
1678	1542	NM_017193	j,I,m,z	II (Kat2), mRNA. Length = 1828	kynurenine aminotransferase II

TABUE 1: E					Ally. Docket No. 44921-51199Wo Doc. No. 1799397.1
Sequence ID No.	ldeniller	GenBank Acel Ref. Seq.(D.	Model Code	Cene Name	واللا بعنعيان ويتوزينا
1679	14695	NM_017202	q,s	Rattus norvegicus cytochrome c oxidase, subunit IVa (Cox4a), mRNA. Length = 696 Rattus norvegicus cytochrome c oxidase,	cytochrome c oxidase, subunit IV
1679	14694	NM_017202	s,z	subunit IVa (Cox4a), mRNA. Length = 696	cytochrome c oxidase, subunit IV
1680	1428	NM_017213	m	Rattus norvegicus outer dense fiber of sperm tails 2 (Odf2), mRNA. Length = 2451	outer dense fiber of sperm tails 2
1681	1622	NM_017216	g.j.s.z	Rattus norvegicus solute carrier family 3, member 1 (Slc3a1), mRNA. Length = 2305 Rattus norvegicus 6-pyruvoyl tetrahydropterin synthase	solute carrier family 3, member 1
1682	13642	NM_017220	v	(Pts), mRNA. Length = 1176 Rattus norvegicus 6-pyruvoyl	
1682	19976	NM_017220	w	tetrahydropterin synthase (Pts), mRNA. Length = 1176 Rattus norvegicus organic cationic transporter-like 1	ESTs
1683	1510	NM_017224_	General	(Orctl1), mRNA. Length = 2227	organic cationic transporter-like 1
1684	1811	NM_017228	j,l,m,z	Rattus norvegicus dentatorubral pallidoluysian atrophy (Drpla), mRNA. Length = 4387 Rattus norvegicus eukaryotic translation elongation factor	dentatorubral pallidoluysian atrophy
1686	17563	NM_017245	a,c,e,q	2 (Eef2), mRNA. Length = 2626	eukaryotic translation elongation factor 2
1687	17502	NM_017248	r	Rattus norvegicus heterogeneous nuclear ribonucleoprotein A1 (Hnrpa1), mRNA. Length = 1696 Rattus norvegicus	heterogeneous nuclear ribonucleoprotein A1
1687	17501	NM_017248	×	heterogeneous nuclear ribonucleoprotein A1 (Hnrpa1), mRNA. Length = 1696	heterogeneous nuclear ribonucleoprotein A1
1688	19	NM_017258	v,General	Rattus norvegicus B-cell translocation gene 1, anti- proliferative (Btg1), mRNA. Length = 1464	B-cell translocation gene 1, anti- proliferative
1689	15300	NM_017259	i,v,cc, General	Rattus norvegicus B-cell translocation gene 2, anti- proliferative (Btg2), mRNA. Length = 2519	B-cell translocation gene 2, anti- proliferative
1689	15301	NM_017259	I,m,v,aa,c c, General	Rattus norvegicus B-cell translocation gene 2, anti- proliferative (Btg2), mRNA. Length = 2519 Rattus norvegicus B-cell	B-cell translocation gene 2, anti- proliferative
1689	15299	NM_017259	I,y,cc, General	translocation gene 2, anti- proliferative (Btg2), mRNA. Length = 2519 Rattus norvegicus protease	B-cell translocation gene 2, anti- proliferative
1690	15224	NM_017264	d	(prosome, macropain) 28 subunit, alpha (Psme1), mRNA. Length = 921	protease (prosome, macropain) 28 subunit, alpha

Table1: S	UMMARY				TATTY , Docket No. 44921-5039W0 Doc. No. 1798897.1
Sequence ID No.	ldenWer	Consonk Acc	Model Model	Gene Name	Unigene Cluster Title
				Rattus norvegicus proteasome (prosome, macropain) subunit, alpha type 3 (Psma3), mRNA.	proteasome (prosome, macropain)
1691	3987	NM_017280	bb	Length = 897 Rattus norvegicus	subunit, alpha type 3
				proteasome (prosome, macropain) subunit, alpha type 4 (Psma4), mRNA.	proteasome (prosome, macropain)
1692	1447	NM_017281	1	Length = 1121 Rattus norvegicus proteasome (prosome,	subunit, alpha type 4
1693	15535	NM_017283	s,bb	macropain) subunit, alpha type 6 (Psma6), mRNA. Length = 932	proteasome (prosome, macropain) subunit, alpha type 6
				Rattus norvegicus ATPase, Ca++ transporting, cardiac muscle, slow twitch 2	ATPage Coul transporting gardies
1694	12349	NM_017290	General	(Atp2a2), mRNA. Length = 5648 Rattus norvegicus calcium	ATPase, Ca++ transporting, cardiac muscle, slow twitch 2
				channel, voltage-dependent, L type, alpha 1D subunit (Cacna1d), mRNA. Length =	calcium channel, voltage-dependent,
1695	15819	NM_017298	р	Rattus norvegicus solute	L type, alpha 1D subunit
4000	02005	NNA 047000	v	carrier family 19 (sodium/hydrogen exchanger), member 1 (Slc19a1), mRNA. Length = 2402	solute carrier family 19 (sodium/hydrogen exchanger), member 1
1696	23825	NM_017299		Rattus norvegicus solute carrier family 19 (sodium/hydrogen exchanger), member 1	solute carrier family 19
1696	23826	NM_017299	v	(Slc19a1), mRNA. Length = 2402	(sodium/hydrogen exchanger), member 1
1697	14003	NM 017305	j,l,m,y,z	Rattus norvegicus glutamate- cysteine ligase , modifier subunit (Gclm), mRNA. Length = 1382	Glutamate-cysteine ligase (gamma- glutamylcysteine synthetase), regulatory
				Rattus norvegicus dodecenoyl-Coenzyme A delta isomerase (3,2 trans- enoyl-Coenyme A	
1698	26109	NM_017306	q,s	isomerase) (DCI), mRNA. Length = 972	EST
				Rattus norvegicus dodecenoyl-Coenzyme A delta isomerase (3,2 trans- enoyl-Coenyme A	Rat mRNA for delta3, delta2-enoyl- CoA isomerase,dodecenoyl-
1698	18687	NM_017306	q,t	isomerase) (DCI), mRNA. Length = 972	Coenzyme A delta isomerase (3,2 trans-enoyl-Coenyme A isomerase)
1699	18142	NM_017314	g,s,aa	Rattus norvegicus ubiquitin C (Ubc), mRNA. Length = 2545 Rattus norvegicus cathepsin	ubiquitin C
1700	1894	NM_017320	t	S (Ctss), mRNA. Length = 1330 Rattus norvegicus	cathepsin S
1701	20809	NM_017326	u	calmodulin (RCM3), mRNA. Length = 1112	calmodulin
				Rattus norvegicus transcriptional repressor CREM (CREM), mRNA.	
1702	355	NM_017334	СС	Length = 436	

table 1: S	UMMARY				Aity, Docket No. 44921-5089W0 Doc. No. 1798897.1
Sequence ID No.	ldentifier	GenBank Ace Ref. Seg ID	Model Code	Gene Name	Unigene Cluster Title
1703	16148	NM_017340	q,s	Rattus norvegicus acyl-coA oxidase (RATACOA1), mRNA. Length = 3741	acyl-coA oxidase
1703	16150	NM_017340	a	Rattus norvegicus acyl-coA oxidase (RATACOA1), mRNA. Length = 3741	acyl-coA oxidase
1704	20849	NM_017343	r,u, General	Rattus norvegicus myosin regulatory light chain (MRLCB), mRNA. Length = 1139	Rat mRNA for myosin regulatory light chain (RLC)
			b,General	Rattus norvegicus myosin regulatory light chain (MRLCB), mRNA. Length = 1139	Rat mRNA for myosin regulatory light chain (RLC)
1704	606	NM_017343	b,General	Rattus norvegicus urinary plasminogen activator receptor 2 (uPAR-2), mRNA. Length = 1272	urinary plasminogen activator receptor 2
1706	1581	NM 017365	General	Rattus norvegicus PDZ and LIM domain 1 (elfin) (Pdlim1), mRNA. Length = 1392	LIM protein
1707	455	NM 019131	x	Rattus norvegicus Tropomyosin 1 (alpha) (Tpm1), mRNA. Length = 1004	Tropomyosin 1 (alpha)
1707		14W_013131		Rattus norvegicus Tropomyosin 1 (alpha) (Tpm1), mRNA. Length =	
1707	456	NM_019131	ly,z	Rattus norvegicus Solute carrier family 12, member 1 (bumetanide-sensitive sodium-[potassium]-chloride cotransporter) (Sic12a1),	Tropomyosin 1 (alpha) Solute carrier family 12, member 1 (bumetanide-sensitive sodium-
1708	4532	NM_019134	b	mRNA. Length = 4595 Rattus norvegicus synaptogyrin 1 (Syngr1),	[potassium]-chloride cotransporter) ESTs, Moderately similar to synaptogyrin
1709	1608	NM_019166	j,y,z	mRNA. Length = 879 Rattus norvegicus synuclein, alpha (Snca), mRNA. Length	[R.norvegicus],synaptogyrin 1
1710	7489	NM_019169	c,General	= 1018 Rattus norvegicus carbonyl reductase (Cbr), mRNA.	synuclein, alpha
1711	17066	NM_019170	p	Rattus norvegicus carbonic anhydrase 4 (Ca4), mRNA.	carbonyl reductase ESTs, Highly similar to CARBONIC ANHYDRASE IV PRECURSOR [R.norvegicus]
1712	23924	NM_019174 NM_019186	bb t	Length = 1205 Rattus norvegicus ADP- ribosylation-like 4 (Arl4), mRNA. Length = 1067	ADP-ribosylation-like 4
1714	22063	NM_019195	d	Rattus norvegicus integrin- associated protein (Cd47), mRNA. Length = 1053	integrin-associated protein
1715	2079	NM_019220	j,k,z	Rattus norvegicus amino- terminal enhancer of split (Aes), mRNA. Length = 1356 Rattus norvegicus solute	related to Drosophila groucho gene
1716	16284	NM_019229	l,m	carrier family 12, member 4 (Slc12a4), mRNA. Length = 3726 Rattus norvegicus small	solute carrier family 12, member 4
1717	985_	NM_019233	b,cc	inducible cytokine subfamily A20 (Scya20), mRNA. Length = 816	small inducible cytokine subfamily A20

TABLE 18 S	WXXXXXXV				///fiy. Docket No. 4/1921-5069W0 Doc. No. 1793397.1
Sequence ID No.		Conbank Acol Roi, Seq ID	Model Code	Cene Name	Uniterne Christer Title
1718	15503	NM 019237	k,x	Rattus norvegicus procollagen C-proteinase enhancer protein (Pcolce), mRNA. Length = 1547	procollagen C-proteinase enhancer protein
				Rattus norvegicus procollagen C-proteinase enhancer protein (Pcolce),	procollagen C-proteinase enhancer protein
1718	15504	NM_019237	k,x	mRNA. Length = 1547 Rattus norvegicus interferon- related developmental regulator 1 (Ifrd1), mRNA.	interferon-related developmental
1719	17908	NM_019242	I,v,cc, General	Length = 1736 Rattus norvegicus paired-like	regulator 1
1720	11218	NM_019247	С	homeodomain transcription factor 3 (Pitx3), mRNA. Length = 1253	paired-like homeodomain transcription factor 3
4704	45250	NIM 010250	d,f	Rattus norvegicus complement component 1, q subcomponent binding protein (C1qbp), mRNA. Length = 1124	complement component 1, q subcomponent binding protein
1721	21443	NM_019259 NM_019262	aa, General	Rattus norvegicus complement component 1, q subcomponent, beta polypeptide (C1qb), mRNA. Length = 1136	complement component 1, q subcomponent, beta polypeptide
4700		NIA 040250	. Constal	Rattus norvegicus complement component 1, q subcomponent, beta polypeptide (C1qb), mRNA.	complement component 1, q
1723	117	NM_019262	t,General	Length = 1136 Rattus norvegicus sodium channel, voltage-gated, type VIII, alpha polypeptide (Scn8a), mRNA. Length = 6586	subcomponent, beta polypeptide sodium channel, voltage-gated, type VIII, alpha polypeptide
4724	1145	NM 019280	w	Rattus norvegicus gap junction membrane channel protein alpha 5 (Gja5), mRNA. Length = 3115	gap junction membrane channel protein alpha 5
1724	22220	NM 019286	С	Rattus norvegicus Alcohol dehydrogenase 3 (Adh3), mRNA. Length = 1131	Alcohol dehydrogenase (class I), alpha polypeptide
1726	10015	NM 019289		Rattus norvegicus Actin- related protein complex 1b (Arpc1b), mRNA. Length = 1430	Actin-related protein complex 1b
1726	10016	NM_019289	bb, General	Rattus norvegicus Actin- related protein complex 1b (Arpc1b), mRNA. Length = 1430	Actin-related protein complex 1b
1727	21651	NM_019296	c,f,x	Rattus norvegicus Cell division cycle control protein 2 (Cdc2a), mRNA. Length = 1184	Cell division cycle control protein 2
1728	20751	NM_019301	s	Rattus norvegicus Complement receptor related protein (Cr1), mRNA. Length = 1811	Complement receptor related protein
1729	645	NM_019345	bb	Rattus norvegicus solute carrier family 12, member 3 (Slc12a3), mRNA. Length = 4361	solute carrier family 12, member 3

TABLE 1: 8					/Aiiy, Dockei No. 44921-5089W0 Doc. No. 1793397.1
Scancinco D No.	ldentifier.	GenBank Ace Ref. Seq ID	Model Code	Gene Name	ese (स्ट. १७००) । Unigene Cluster Tille
1730	1301	NM_019349	c	Rattus norvegicus Serine/threonine kinase 2 (Stk2), mRNA. Length = 4194	Rat liver stearyl-CoA desaturase mRNA, complete cds
1731	3776	NM_019354	a,u	Rattus norvegicus Uncoupling protein 2, mitochondrial (Ucp2), mRNA. Length = 1575 Rattus norvegicus eukaryotic	Uncoupling protein 2, mitochondrial
1732	4592	NM_019356	General	translation initiation factor 2, subunit 1 (alpha) (Eif2s1), mRNA. Length = 1377 Rattus norvegicus factor-	eukaryotic translation initiation factor 2, subunit 1 (alpha)
1733	1324	NM_019371	w	responsive smooth muscle protein (SM-20), mRNA. Length = 2825 Rattus norvegicus 14-3-3	factor-responsive smooth muscle protein
1734	19577	NM_019377	e	protein beta-subtype (Ywhab), mRNA. Length = 2756 Rattus norvegicus Testis	ESTs, Moderately similar to S12207 hypothetical protein [M.musculus]
1735	24626	NM_019381	s	enhanced gene transcript (Tegt), mRNA. Length = 940 Rattus norvegicus espin (Espn), mRNA. Length =	Testis enhanced gene transcript
1736	744	NM_019622	р	2786 Rattus norvegicus cytochrome P450 4F1	espin
1737	20716	NM_019623	С	(Cyp4f1), mRNA. Length = 1977 Rattus norvegicus beta-galactoside-binding lectin	cytochrome P450 4F1
1738	20709	NM_019904	x	(Lgals1), mRNA. Length = 519	beta-galactoside-binding lectin Rattus norvegicus clone BB.1.4.1
1739	574	NM_019905	u,General		unknown Glu-Pro dipeptide repeat protein mRNA, complete cds,calpactin I heavy chain,hydroxyacid oxidase 3 (medium-chain)
1740	9096	NM_019908	j	hypothetical protein LOC56728 (LOC56728), mRNA. Length = 858	hypothetical protein LOC56728
1741	20457	NM 020073	i,General	Rattus norvegicus parathyroid hormone receptor (LOC56813), mRNA. Length = 2065	parathyroid hormone receptor
1741	20458	NM_020073	General	Rattus norvegicus parathyroid hormone receptor (LOC56813), mRNA. Length = 2065	parathyroid hormone receptor
1741	20460	NM_020073	General	Rattus norvegicus parathyroid hormone receptor (LOC56813), mRNA. Length = 2065	parathyroid hormone receptor
1742	18713	NM_020075	r	Rattus norvegicus eukaryotic initiation factor 5 (eIF-5) (Eif5), mRNA. Length = 3504	eukaryotic initiation factor 5 (eIF-5)
1742	18715	NM_020075	r	Rattus norvegicus eukaryotic initiation factor 5 (eIF-5) (Eif5), mRNA. Length = 3504	eukaryotic initiation factor 5 (eIF-5)

	SUMMARY	7g - 2			*Ally, Docket No. 44921-503900 Doc. No. 1793397,
Sequence ID No.	(Identifier	Generik Ace/ Ref. Sec ID	Model Code	Gene Name	Unigene Civster Title
				Rattus norvegicus 3- hydroxyanthranilate 3,4- dioxygenase (Haao), mRNA.	3-hydroxyanthranilate 3,4-
1743	20493	NM_020076	р	Length = 1254 Rattus norvegicus kidney- specific membrane protein	dioxygenase
1744	16375	NM_020976	g	(NX-17), mRNA. Length = 1181 Rattus norvegicus thymosin,	kidney-specific membrane protein
1745	20816	NM_021261	k,General	beta 10 (Tmsb10), mRNA. Length = 539 Rattus norvegicus ribosomal	thymosin beta-10
1746	15335	NM_021264	a	protein L35a (Rpl35), mRNA. Length = 348 Rattus norvegicus	ribosomal protein L35a
1747	18729	NM_021578	k,z	transforming growth factor beta-1 gene (Tgfb1), mRNA. Length = 1585	transforming growth factor beta-1 gene
				Rattus norvegicus transforming growth factor- beta (TGF-beta) masking protein large subunit (Ltbp1),	transforming growth factor-beta (TGF
1748	19060	NM_021587	сс	mRNA. Length = 6244 Rattus norvegicus	beta) masking protein large subunit
1749	17324	NM_021593	o,General	kynurenine 3-hydroxylase (Kmo), mRNA. Length = 1733 Rattus norvegicus Thyroxine	kynurenine 3-hydroxylase
1750	19679	NM_021653	General	deiodinase, type I (Dio1), mRNA. Length = 2106 Rattus norvegicus Thyroxine	Thyroxine deiodinase, type I
1750	19678	NM_021653	a,v, General	deiodinase, type I (Dio1), mRNA. Length = 2106 Rattus norvegicus putative	Thyroxine delodinase, type I
1751	19665	NM_021688	u,General	potassium channel TWIK (Kcnk1), mRNA. Length = 1582	putative potassium channel TWIK
				Rattus norvegicus cAMP- regulated guanine nucleotide exchange factor I (cAMP-	
1752	19667	NM_021690	m	GEFI) (Epac), mRNA. Length = 3373 Rattus norvegicus	cAMP-regulated guanine nucleotide exchange factor I (cAMP-GEFI)
1754	22916	NM_021740	a	prothymosin alpha (Ptma), mRNA. Length = 1182 Rattus norvegicus CD14	prothymosin alpha
1755	19710	NM_021744	t	antigen (Cd14), mRNA. Length = 1591 Rattus norvegicus CD14	CD14 antigen
1755	19711	NM_021744	t	antigen (Cd14), mRNA. Length = 1591 Rattus norvegicus farnesoid	CD14 antigen
1756	19712	NM_021745	r	X activated receptor (LOC60351), mRNA. Length = 2070	farnesoid X activated receptor
1757	1962	NIM 021750	ikva	Rattus norvegicus cysteine- sulfinate decarboxylase (Csad), mRNA. Length = 2413	Rattus norvegicus cca2 mRNA,
1757	1902	NM_021750	j,k,y,z	Rattus norvegicus cysteine- sulfinate decarboxylase (Csad), mRNA. Length =	complete cds
1757	19824	NM_021750	a,bb	2413 Rattus norvegicus Nopp140 associated protein (Nap65),	cysteine-sulfinate decarboxylase
1758	25198	NM_021754	h	mRNA. Length = 1980	Nopp140 associated protein

TABLE 18' S	WIMMARY				Alty, Docket No. 44921-5089WO Doc. No. 1793897.1
Sequence ID No.	 ຄ່າວຄານໃ <u>ງ</u> ກະ	GenBank Acc/ Ref. Sec ID	Model Gode	©ano Namo	Unitene Greter Tille
1758	20035	NM_021754	b,n,s,v, General	Rattus norvegicus Nopp140 associated protein (Nap65), mRNA. Length = 1980	Nopp140 associated protein
1759	20090	NM_021757	m	Rattus norvegicus pleiotropic regulator 1 (Plrg1), mRNA. Length = 1545	pleiotropic regulator 1
1760	17885	NM_021765	aa	Rattus norvegicus beta prime COP (Copb), mRNA. Length = 3025	beta prime COP
1762	20161	NM_021836	cc, General	Rattus norvegicus jun B proto-oncogene (Junb), mRNA. Length = 1035	jun B proto-oncogene
1764	1203	NM_021997	k,z	Rattus norvegicus cytoplasmic linker 2 (Cyln2), mRNA. Length = 4847	cytoplasmic linker 2
1765	23151	NM_022005	b	mRNA. Length = 1711	FXYD domain-containing ion transport regulator 6
1767	17101	NM_022179	bb	Rattus norvegicus Hexokinase 3 (Hk3), mRNA. Length = 3692	Hexokinase 3
1767	17100	NM_022179	рр	Rattus norvegicus Hexokinase 3 (Hk3), mRNA. Length = 3692	Hexokinase 3
1768	20257	NM_022180	w, General	Rattus norvegicus Hepatic nuclear factor 4(alpha transcription factor 4) (Hnf4a), mRNA. Length = 1446	Hepatic nuclear factor 4 (alpha transcription factor 4)
				Rattus norvegicus Hepatic nuclear factor 4(alpha transcription factor 4) (Hnf4a), mRNA. Length =	Hepatic nuclear factor 4 (alpha
1768	25699	NM_022180	İ	1446 Rattus norvegicus Hepatic nuclear factor 4(alpha transcription factor 4) (Hnf4a), mRNA. Length =	transcription factor 4)
1768	10860	NM_022180	р	Rattus norvegicus topoisomerase (DNA) II	ESTs
1769	23780	NM_022183	k,x	alpha (Top2a), mRNA. Length = 6052 Rattus norvegicus resiniferatoxin-binding.	topoisomerase (DNA) II alpha
1770	20312	NM_022224	0	phosphotriesterase-related protein (Rpr1), mRNA. Length = 1050 Rattus norvegicus connective tissue growth	resiniferatoxin-binding, phosphotriesterase-related protein
1771	6585	NM_022266	d,p,cc	factor (Ctgf), mRNA. Length = 2345 Rattus norvegicus alpha-	connective tissue growth factor
1772	17161	NM_022298	i,v,cc, General	tubulin (Tuba1), mRNA. Length = 1617 Rattus norvegicus alpha-	alpha-tubulin
1772	17162	NM_022298	u	tubulin (Tuba1), mRNA. Length = 1617 Rattus norvegicus alpha-	alpha-tubulin
1772	17160	NM_022298	u	tubulin (Tuba1), mRNA. Length = 1617 Rattus norvegicus alpha-	alpha-tubulin
1772	17158	NM_022298	q .	tubulin (Tuba1), mRNA. Length = 1617	alpha-tubulin

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TABLE 1: 8	BUMMARY				Atty, Doctot No. 44921-5089W0 Doc. No. 1793897.1
Echence Sechence	ldentitier	GenBenk Acc Ref. Seg ID	Model Code	Cene Name	Unigeno Cluster Ville *
			i,aa,	Rattus norvegicus Proliferating cell nuclear antigen (Pcna), mRNA.	
1773	11454	NM_022381	General	Length = 1160 Rattus norvegicus	Proliferating cell nuclear antigen
1773	11455	NM_022381	I,General	Proliferating cell nuclear antigen (Pcna), mRNA. Length = 1160	Proliferating cell nuclear antigen
				Rattus norvegicus quinoid dihydropteridine reductase (Qdpr), mRNA. Length =	
1774	13480	NM_022390	s	1307	quinoid dihydropteridine reductase
1775	15184	NM_022391	z	Rattus norvegicus pituitary tumor-transforming 1 (Pttg1), mRNA. Length = 974	pituitary tumor transforming gene
				Rattus norvegicus growth response protein (CL-6) (LOC64194), mRNA. Length	
1776	22413	NM_022392	h	= 2410 Rattus norvegicus growth	growth response protein (CL-6)
1776	22414	NM 022392	n	response protein (CL-6) (LOC64194), mRNA. Length = 2410	growth response protein (CL-6)
				Rattus norvegicus macrophage galactose N- acetyl-galactosamine specific lectin (MgI), mRNA.	
1777	22499	NM_022393	t	Length = 1358	Gal/GalNAc-specific lectin
				Rattus norvegicus calreticulin (Calr), mRNA.	
1779	24537	NM_022399	е	Length = 1882 Rattus norvegicus	calreticulin
				calreticulin (Calr), mRNA.	
1779	24539	NM_022399	У	Length = 1882 Rattus norvegicus plectin	calreticulin
4700		NIN 000404	o,General	(Plec1), mRNA. Length =	nlostin
1780	1141	NM_022401	o,General	Rattus norvegicus acidic	plectin
1781	1069	NM 022402	g	ribosomal protein P0 (Arbp), mRNA. Length = 1046	acidic ribosomal protein P0
1782	8211	NM 022500	j,n,s	Rattus norvegicus ferritin light chain 1 (FtI1), mRNA. Length = 552	ferritin light chain 1
1782	8212	NM 022500	n,s	Rattus norvegicus ferritin light chain 1 (Ftl1), mRNA. Length = 552	ferritin light chain 1
1762	0212	14W_022300	11,5	Rattus norvegicus cytochrome c oxidase	Territin light chair i
1783	6815	NM_022503	s	subunit VIIa 3 (Cox7a3), mRNA. Length = 460 Rattus norvegicus ribosomal	cytochrome c oxidase subunit VIIa 3
1784	4259	NM_022504	q,w	protein L36 (Rpl36), mRNA. Length = 364	ribosomal protein L36
				Rattus norvegicus survival motor neuron (Smn), mRNA.	
1785	1611	NM_022509	<u>ii</u>	Length = 1243	survival motor neuron
				Rattus norvegicus short chain acyl-coenzyme A	
1786	2236	NM_022512	y,z	dehydrogenase (Acads), mRNA. Length = 1749 Rattus norvegicus ribosomal	short chain acyl-coenzyme A dehydrogenase
1787	3026	NM_022514	а	protein L27 (Rpl27), mRNA. Length = 463	ribosomal protein L27

TABLE 1: 8	BUIMMARY				Ally. Docket No. 44221-5639W0 Doc. No. 1793897.1
Sequence ID No.	ldeniiier	Genfank Acc Ref. Seg ID	Model Model	Cene Namo	Unigene Custer Tille
1787	3027	NM_022514	a,q,r,aa	Rattus norvegicus ribosomal protein L27 (Rpl27), mRNA. Length = 463	ribosomal protein L27
1788	2696	NM_022515	a,d	Rattus norvegicus ribosomal protein L24 (Rpl24), mRNA. Length = 541	ribosomal protein L24
1788	2697	NM_022515	n,w,aa	Rattus norvegicus ribosomal protein L24 (Rpl24), mRNA. Length = 541 Rattus norvegicus	ribosomal protein L24
1789	3900	NM_022516	h	polypyrimidine tract binding protein (Ptb), mRNA. Length = 2697 Rattus norvegicus ADP-	polypyrimidine tract binding protein
1790	4151	NM_022518	o	ribosylation factor 1 (Arf1), mRNA. Length = 900 Rattus norvegicus ornithine	ADP-ribosylation factor 1
1791	4242	NM_022521	С	aminotransferase (Oat), mRNA. Length = 1938 Rattus norvegicus platelet	ornithine aminotransferase
1792	4412	NM_022523	0	endothelial tetraspan antigen 3 (Cd151), mRNA. Length = 1668	platelet endothelial tetraspan antigen- 3
1793	6641	NM_022533	General	Rattus norvegicus plasmolipin (Z49858), mRNA. Length = 1475 Rattus norvegicus cyclophilin	plasmolipin
1794	8097	NM_022536	а	B (Ppib), mRNA. Length = 840 Rattus norvegicus	cyclophilin B
1795	8597	NM 022538	c,r,u	phosphatidate phosphohydrolase type 2 ((Ppap2), mRNA. Length =	phosphatidate phosphohydrolase type 2
1795	8598	NM 022538	u	Rattus norvegicus phosphatidate phosphohydrolase type 2 (Ppap2), mRNA. Length = 871	phosphatidate phosphohydrolase type
1735	0030	14W_022330		Rattus norvegicus small zinc finger-like protein DDP2	
1796	9296	NM_022541	0	(Ddp2), mRNA. Length = 494 Rattus norvegicus ornithine decarboxylase antizyme	small zinc finger-like protein DDP2
1797	21063	NM_022585	h	inhibitor (Oazi), mRNA. Length = 4269 Rattus norvegicus	ornithine decarboxylase antizyme inhibitor
1799	20781	NM_022591	z	telomerase protein component 1 (Tlp1), mRNA. Length = 8216	telomerase protein component 1
1800	20803	NM_022592	n	Rattus norvegicus transketolase (Tkt), mRNA. Length = 2098 Rattus norvegicus enoyl	transketolase
1801	20925	NM_022594	q	hydratase-like protein, peroxisomal (Ech1), mRNA. Length = 1097	enoyl hydratase-like protein, peroxisomal
1802	20944	NM_022597	aa	Rattus norvegicus cathepsin B (Ctsb), mRNA. Length = 1904	cathepsin B
1803	21024	NM_022599	o,General	Rattus norvegicus synaptojanin 2 binding protein (Synj2bp), mRNA. Length = 5215	outer membrane protein

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UABLE 18					Aug, Docket No. 44921-5099W0 Doc. No. 1793997.1
Sequence ID No.	ldentiller	Conedix Acci Rei. Seq ID	Model Code	Cene Name	Unigene Civeta; Tille
1804	2250	NM_022643	General	Rattus norvegicus Testis- specific histone 2b (Th2b), mRNA. Length = 470	ESTs, Highly similar to 0506206A histone H2B [R.norvegicus]
1805	17567	NM_022672	а,у	Rattus norvegicus ribosomal protein S14 (Rps14), mRNA. Length = 492	ribosomal protein S14
4900	17661	NINA 022674	hh	Rattus norvegicus H2A histone family, member Z (H2afz), mRNA. Length = 811	H2A history family, member 7
1806	17661	NM_022674	bb	Rattus norvegicus protein phosphatase 1, regulatory (inhibitor) subunit 1A (Ppp1r1a), mRNA. Length =	H2A histone family, member Z protein phosphatase 1, regulatory
1807	24563	NM_022676	b	Rattus norvegicus protein phosphatase 1, regulatory ((inhibitor) subunit 1A	(inhibitor) subunit 1A
1807	24564	NM_022676	b,x	(Ppp1r1a), mRNA. Length = 619 Rattus norvegicus germinal	protein phosphatase 1, regulatory (inhibitor) subunit 1A
1808	20506	NM_022686	1	histone H4 gene (Hist4), mRNA. Length = 377 Rattus norvegicus preoptic	germinal histone H4 gene
1809	20508	NM_022688	g	regulatory factor-1 (Porf1), mRNA. Length = 689 Rattus norvegicus p105	preoptic regulatory factor-1
1810	17586	NM_022694	k	coactivator (U83883), mRNA. Length = 3166 Rattus norvegicus ribosomal	p105 coactivator
1811	17730	NM_022697	а	protein L28 (Rpl28), mRNA. Length = 466 Rattus norvegicus ribosomal	ribosomal protein L28
1811	17729	NM_022697	q	protein L28 (Rpl28), mRNA. Length = 466 Rattus norvegicus crp-ductin	ribosomal protein L28
1812	154	NM_022849	t	(Crpd), mRNA. Length = 4344 Rattus norvegicus casein	crp-ductin
1813	127	NM_022855	h	kinase 1 gamma 3 isoform (Csnk1g3), mRNA. Length = 2547	casein kinase 1 gamma 3 isoform
1814	152	NM_022858	j	Rattus norvegicus HNF- 3/forkhead homolog-1 (Hfh1), mRNA. Length = 1760	HNF-3/forkhead homolog-1
1816	18101	NM_022948	z	Rattus norvegicus tricarboxylate carrier-like protein (Loc65042), mRNA. Length = 2699	tricarboxylate carrier-like protein
	49400	NNA 000040		Rattus norvegicus tricarboxylate carrier-like protein (Loc65042), mRNA.	trion thousand a coming like
1816	18103	NM_022948	u	Length = 2699 Rattus norvegicus putative protein phosphatase 1 nuclear targeting subunit (Ppp1r10), mRNA. Length =	tricarboxylate carrier-like protein
1817	21491	NM_022951	w	4131 Rattus norvegicus phosphatidylinositol 3-kinase	nuclear targeting subunit
1818	15742	NM_022958	у	(Pik3c3), mRNA. Length = 2752	phosphatidylinositol 3-kinase

TABLE 18 8	BUMMARY				Ally, Docket No. 44221-5032W0 Doc. No. 1793327.1
Sequence ID No.	ldenmer	GenBank Acci. Ref. Seq ID	Model Code	Cene Name	Unigene Givster Tible
1819	9286	NM_023027	t,w	Rattus norvegicus tRNA selenocysteine associated protein (Secp43), mRNA. Length = 864	tRNA selenocysteine associated protein
1820	23215	NM_023102	z	Rattus norvegicus casein kinase 1 gamma 2 isoform (Csnk1g2), mRNA. Length = 1572	casein kinase 1 gamma 2 isoform
1821	21238	NM 024125	cc, General	Rattus norvegicus Liver activating protein (LAP,also NF-IL6, nuclear factor-IL6, previously designated TCF5) (Cebpb), mRNA. Length = 1408	Liver activating protein (LAP, also NF- IL6, nuclear factor-IL6, previously designated TCF5)
1821	21239	NM_024125	cc, General	Rattus norvegicus Liver activating protein (LAP,also NF-IL6, nuclear factor-IL6, previously designated TCF5) (Cebpb), mRNA. Length = 1408	Liver activating protein (LAP, also NF- IL6, nuclear factor-IL6, previously designated TCF5)
1822	353	NM_024127	i,n, General	Rattus norvegicus DNA- damage-inducible transcript 1 (Gadd45a), mRNA. Length = 711	DNA-damage-inducible transcript 1
1822	354	NM_024127	i,n, General	Rattus norvegicus DNA- damage-inducible transcript 1 (Gadd45a), mRNA. Length = 711	DNA-damage-inducible transcript 1
1822	352	NM_024127	h,General	Rattus norvegicus DNA- damage-inducible transcript 1 (Gadd45a), mRNA. Length = 711	DNA-damage-inducible transcript 1
1823	17227	NM_024131	x	Rattus norvegicus D- dopachrome tautomerase (Ddt), mRNA. Length = 628 Rattus norvegicus DNA-	D-dopachrome tautomerase
1824	1598	NM_024134	1	damage inducible transcript 3 (Ddit3), mRNA. Length = 806 Rattus norvegicus	DNA-damage inducible transcript 3
1825	1162	NM_024153	d	adrenodoxin reductase (Fdxr), mRNA. Length = 1786	adrenodoxin reductase
1826	7863	NM_024156	С	Rattus norvegicus annexin VI (Anxa6), mRNA. Length = 2739 Rattus norvegicus	Rattus norvegicus mRNA for H(+)- transporting ATPase, complete cds
1827	22079	NM_024157	x	complement factor I (Cfi), mRNA. Length = 2021	complement factor I
1828	16476	NM_024162	General	Rattus norvegicus heart fatty acid binding protein (Fabp3), mRNA. Length = 666 Rattus norvegicus heat shock 70kD protein 8	heart fatty acid binding protein
1829	17765	NM_024351	b,s,v	(Hspa8), mRNA. Length = 2073 Rattus norvegicus hairy and enhancer of split 1,	Heat shock cognate protein 70
1830	8879	NM_024360	h	(Drosophila) (Hes1), mRNA. Length = 1453	hairy and enhancer of split 1, (Drosophila)

TABLE 1: S	OMMARY			Šitu	Ally, Docket No. 44921-5039W0 Doc. No. 1793397.1
Sequence ID No.	ldentifier.	Cenbenk Ace/ Ref. Seq ID	Model Gode	Come Name	Unigene Cluster Tillo
1831	20772	NM 024363	x	Rattus norvegicus heterogeneous nuclear ribonucleoproteins methyltransferase-like 2 (S. cerevisiae) (Hrmt112), mRNA. Length = 1201	heterogeneous nuclear ribonucleoproteins methyltransferase- like 2 (S. cerevisiae)
1832	2812	NM 024386	С	Rattus norvegicus 3-hydroxy- 3-methylglutaryl CoA lyase (Hmgcl), mRNA. Length = 1390	3-hydroxy-3-methylglutaryl CoA lyase
1833	335	NM 024387	j,y	Rattus norvegicus heme oxygenase-2 non-reducing isoform (Hmox2), mRNA. Length = 1815	heme oxygenase-2 non-reducing isoform
1834	21	NM_024388	cc	Rattus norvegicus immediate early gene transcription factor NGFI-B (Nr4a1), mRNA. Length = 2488	
1834	22	NM_024388	СС	Rattus norvegicus immediate early gene transcription factor NGFI-B (Nr4a1), mRNA. Length = 2488	immediate early gene transcription factor NGFI-B
1836	9929	NM_024392	f	Rattus norvegicus peroxisomal multifunctional enzyme type II (Hsd17b4), mRNA. Length = 2535	peroxisomal multifunctional enzyme type II
1837	3582	NM 024396	aa	Rattus norvegicus ATP- binding cassette, sub-family A (ABC1), member 2 (Abca2), mRNA. Length = 8040	ATP-binding cassette, sub-family A (ABC1), member 2
1838	19993	NM_024398	e,p,s,aa	Rattus norvegicus mitochondrial aconitase (nuclear aco2 gene) (Aco2), mRNA. Length = 2744	mitochondrial aconitase (nuclear aco2 gene)
1839	10789	NM_024399	o	Rattus norvegicus aspartoacylase (Aspa), mRNA. Length = 1552 Rattus norvegicus a	aspartoacylase
1840	22626	NM_024400	cc, General	disintegrin and metalloproteinase with thrombospondin motifs 1 (ADAMTS-1) (Adamts1), mRNA. Length = 4878	a disintegrin and metalloproteinase with thrombospondin motifs 1 (ADAMTS-1)
1841	13633	NM_024403	o General	Rattus norvegicus activating transcription factor ATF-4 (Atf4) mRNA Length = 1173	activating transcription factor ATF-4
1841	13634	NM 024403		Rattus norvegicus activating transcription factor ATF-4	activating transcription factor ATF-4
1842	23387	NM_024404	b,General	Rattus norvegicus RNA binding protein p45AUF1 (Hnrpd), mRNA. Length = 1240	RNA binding protein p45AUF1
1843	21038	NM_024484	h	Rattus norvegicus aminolevulinic acid synthase 1 (Alas1), mRNA. Length = 2052 Rattus norvegicus	aminolevulinic acid synthase 1
1844	1853	NM_030826	s	Glutathione peroxidase 1 (Gpx1), mRNA. Length = 1539	ESTs,Glutathione peroxidase 1

TABLE1:	SUMMARY				Ally, Docket No. 44921-5039W0 Doc. No. 1793397.1
Scalience ID No:	ldenWier	Conbank Acc/ Ref. Scq ID	Model Code	Gene Name	Unicene Greater Tille
1845	15111	NM_030827	e,General	Rattus norvegicus glycoprotein 330 (Lrp2), mRNA. Length = 15,438	glycoprotein 330
1845	15112	NM_030827	y,z	Rattus norvegicus glycoprotein 330 (Lrp2), mRNA. Length = 15,438	glycoprotein 330
1845	15110	NM_030827	General	Rattus norvegicus glycoprotein 330 (Lrp2), mRNA. Length = 15,438	glycoprotein 330
1846	808	NM 030837	k,m	Rattus norvegicus kidney specific organic anion transporter (Slc21a4), mRNA. Length = 2772	kidney specific organic anion transporter
1646	808	NNI_030637	K,III	Rattus norvegicus islet cell autoantigen 1, 69 kDa (Ica1),	mansporter
1847	4057	NM_030844	k	mRNA. Length = 2094	islet cell autoantigen 1, 69 kDa
	1			Rattus norvegicus gro	
1848	1221	NM_030845	t	(Gro1), mRNA. Length = 929 Rattus norvegicus epithelial	gro
				membrane protein 3 (Emp3),	
1849	21509	NM_030847	x	mRNA. Length = 737	epithelial membrane protein 3
				Rattus norvegicus pyruvate dehydrogenase kinase 2	
				subunit p45 (PDK2) (Pdk2),	pyruvate dehydrogenase kinase 2
1850	1928	NM_030872	v	mRNA. Length = 2207	subunit p45 (PDK2)
				Rattus norvegicus profilin II	
1851	17342	NM 030873	u	(Pfn2), mRNA. Length = 1966	profilin II
1001	17342	14141_00007.5		Rattus norvegicus	promit ii
				Angiotensin II receptor, type	
1852	24648	NM 030985	u	1 (AT1A) (Agtr1a), mRNA. Length = 1450	Angiotensin II receptor, type 1 (AT1A)
1032	24048	14141_030903	u -	Rattus norvegicus	Angiotensii ii receptor, type ii (Ai iA)
				Angiotensin II receptor, type	
4050	25452	NM 020095	Conorni	1 (AT1A) (Agtr1a), mRNA. Length = 1450	
1852	25453	NM_030985	General	Rattus norvegicus Guanine	
				nucleotide-binding protein	
				beta 1 (Gnb1), mRNA.	Guanine nucleotide-binding protein
1853	21802	NM_030987	h	Length = 2837 Rattus norvegicus aldo-keto	beta 1
				reductase family 1, member	
				A1 (aldehyde reductase)	
1854	23109	NM 031000	f,s,z	(Akr1a1), mRNA. Length =	aldo-keto reductase family 1, member A1 (aldehyde reductase)
1004	20103	1.1.1001000	.,5,2	Rattus norvegicus 4-	(Liberty de l'addition)
				aminobutyrate	
1855	134	NM 031003	a,u	aminotransferase (Abat), mRNA. Length = 1726	4-aminobutyrate aminotransferase
,300	104	1414_001000	u,u	Rattus norvegicus	- aloodytato aminoranoiciase
				angiotensin II type-1	
1856	25461	NM 031009	0	receptor (Agtr1), mRNA. Length = 2156	angiotensin II type-1 receptor
1000	120701		<u> </u>	Rattus norvegicus	angiotolion ii typo i receptor
		1		arachidonate 12-	
1857	1845	NM_031010	t	lipoxygenase (Alox12), mRNA. Length = 2048	arachidonate 12-lipoxygenase
1337	1043	11111_001010	-	Rattus norvegicus	and made in a sport goridad
				arachidonate 12-	
1857	25517	NM 031010	c,t	lipoxygenase (Alox12), mRNA. Length = 2048	arachidonate 12-lipoxygenase
1001	2331/	14IVI_031010	, t	Rattus norvegicus p38	aracindonate 12-lipoxygenase
				mitogen activated protein	
1050	16560	NIM 024000	,	kinase (Mapk14), mRNA.	n38 mitagen activated protein kinese
1858	16562	NM_031020	[f	Length = 3132	p38 mitogen activated protein kinase

TABLE 1: S	UMMARY				Ally. Docket No. 44921-5039W0
Seguence		GenBank Acc/	Model		<u>1008. (No. 1793897.1</u>
Sequence ID No.	ldeniiiler	Ref. Seq ID	මානම	Gene Name	Unigene Cluster Tille
1859	1480	NM 031021	f	Rattus norvegicus casein kinase II beta subunit (Csnk2b), mRNA. Length = 1964	casein kinase II beta subunit
1860	1719	NM 031024	n	Rattus norvegicus drebrin A (Dbn1), mRNA. Length = 2697	drebrin A
1861	1350	NM 031030	h	Rattus norvegicus cyclin G- associated kinase (Gak), mRNA. Length = 4454	cyclin G-associated kinase
1862	16775	NM_031031	General	Rattus norvegicus L- arginine: glycine amidinotransferase (Gatm), mRNA. Length = 2260	L-arginine: glycine amidinotransferase
1863	691	NM_031034	w	Rattus norvegicus guanine nucleotide binding protein (G protein) alpha 12 (Gna12), mRNA. Length = 1423	guanine nucleotide binding protein (G protein) alpha 12
1864	15886	NM_031035	z	Rattus norvegicus GTP- binding protein (G-alpha-i2) (Gnai2), mRNA. Length = 1748	GTP-binding protein (G-alpha-i2)
1866	3608	NM_031044	k,General	Rattus norvegicus histamine N-methyltransferase (Hnmt), mRNA. Length = 1225 Rattus norvegicus histamine	histamine N-methyltransferase
1866	3610	NM_031044	d,General	N-methyltransferase (Hnmt), mRNA. Length = 1225 Rattus norvegicus	histamine N-methyltransferase
1867	15137	NM_031051	s	macrophage migration inhibitory factor (Mif), mRNA. Length = 551	macrophage migration inhibitory factor
1868	514	NM 031056	General	Rattus norvegicus matrix metalloproteinase 14, membrane-inserted (Mmp14), mRNA. Length = 2448	matrix metalloproteinase 14, membrane-inserted
1869	17269	NM 031057	General	Rattus norvegicus methylmalonate semialdehyde dehydrogenase gene (Mmsdh), mRNA. Length = 2059	methylmalonate semialdehyde dehydrogenase gene
1870	11849	NM 031065	a	Rattus norvegicus ribosomal protein L10a (Rpl10a), mRNA. Length = 710	ribosomal protein L10a
1871	1855	NM 031074	h	Rattus norvegicus nucleoporin 98 (Nup98), mRNA. Length = 3237	nucleoporin 98
1872	4683	NM_031083	d	Rattus norvegicus phosphatidylinositol 4-kinase (Pik4cb), mRNA. Length = 3205	phosphatidylinositol 4-kinase
1873	15202	NM 031093	а	Rattus norvegicus -ral simian leukemia viral oncogene homolog A (ras related) (Rala), mRNA. Length = 952	#NAME?
		001000		Rattus norvegicus -ral simian leukemia viral oncogene homolog A (ras related)	
1873	15201	NM_031093	a,n	, , ,	#NAME?
1874	12639	NM_031099	aa	Length = 1069	ribosomal protein L5

TABLE 1:	SUMMARY				Ally, Docket No. 44921-5089W0 Doc. No. 1793397.1
Sequence ID No.	ldemijer	Consonk Ace Ref. Seq ID	Model Gode	Cene Name	Unitene Cluster Tille
1875	20812	NM_031100	а	Rattus norvegicus ribosomal protein L10 (Rpl10), mRNA. Length = 769	ribosomal protein L10
1876	16938	NM_031103	w	Rattus norvegicus ribosomal protein L19 (Rpl19), mRNA. Length = 703	ribosomal protein L19
1877	19268	NM_031104	q	Rattus norvegicus ribosomal protein L22 (Rpl22), mRNA. Length = 465	ribosomal protein L22
1878	16929	NM_031108	q	Rattus norvegicus mRNA for ribosomal protein S9 (Rps9), mRNA. Length = 688	mRNA for ribosomal protein S9
1879	10878	NM_031110	q,bb	Rattus norvegicus ribosomal protein S11 (Rps11), mRNA. Length = 534	ribosomal protein S11
1880	19162	NM_031111	aa	Rattus norvegicus ribosomal protein S21 (Rps21), mRNA. Length = 359	ribosomal protein S21
1880	19161	NM_031111	a,bb	Rattus norvegicus ribosomal protein S21 (Rps21), mRNA. Length = 359	ribosomal protein S21
1881	24615	NM_031112	a,y	Rattus norvegicus ribosomal protein S24 (Rps24), mRNA. Length = 466	ribosomal protein S24
1882	20839	NM_031113	a,q	Rattus norvegicus ribosomal protein S27a (Rps27a), mRNA. Length = 552 Rattus norvegicus S-100	ribosomal protein S27a
1883	19040	NM 031114	I,m, General	related protein, clone 42C (S100A10), mRNA. Length = 573	S-100 related protein, clone 42C
				Rattus norvegicus secretin receptor (Sctr), mRNA. Length = 1796	
1884	16349	NM_031115	u	Rattus norvegicus sulfite oxidase (Suox), mRNA.	secretin receptor
1885	14970	NM_031127	General	Length = 1777 Rattus norvegicus thyroid hormone receptor alpha (Thra1), mRNA. Length =	sulfite oxidase
1886	1814	NM_031134	n,q	Rattus norvegicus TGFB inducible early growth	thyroid hormone receptor
1887	13359	NM_031135	General	response (Tieg), mRNA. Length = 3115 Rattus norvegicus thymosin	TGFB inducible early growth response
1888	15052	NM_031136	а	beta-4 (Tmsb4x), mRNA. Length = 686 Rattus norvegicus thymosin	thymosin beta-4
1888	19359	NM_031136	а	beta-4 (Tmsb4x), mRNA. Length = 686	EST
1889	15185	NM_031140	General	Rattus norvegicus vimentin (Vim), mRNA. Length = 1796 Rattus norvegicus	vimentin
1890	21625	NM_031144	a,e	cytoplasmic beta-actin (Actx), mRNA. Length = 1128	cytoplasmic beta-actin
1904	229	NIM 024450	bb	Rattus norvegicus RAB11a, member RAS oncogene family (Rab11a), mRNA.	RAB11a, member RAS oncogene
1891	238	NM_031152 NM_031152	bb	Length = 895 Rattus norvegicus RAB11a, member RAS oncogene family (Rab11a), mRNA. Length = 895	family RAB11a, member RAS oncogene family

TABLE 1: S	UMMARY				Ally, Docket No. 44921-5089WG Doc. No. 1798897.1
Sequence ID No.	(Mentifier	GenBank Acci. Ref. Seq ID.	Modal Code	Gene Name	Unigene Gluster Title
				Rattus norvegicus ubiquitin- conjugating enzyme E2D 3 (homologous to yeast	
1892	15277	NM_031237	g	UBC4/5) (Ube2d3), mRNA. Length = 1531 Rattus norvegicus acyl-CoA	ubiquitin-conjugating enzyme E2D 3 (homologous to yeast UBC4/5)
1893	18083	NM_031315	q	thioesterase 1, cytosolic (Cte1), mRNA. Length =	R.norvegicus mRNA for mitochondrial very-long-chain acyl-CoA thioesterase
				Rattus norvegicus acyl-CoA thioesterase 1, cytosolic (Cte1), mRNA. Length =	R.norvegicus mRNA for mitochondrial very-long-chain acyl-CoA thioesterase,acyl-CoA thioesterase 1,
1893	1858	NM_031315	q	Rattus norvegicus t-complex	cytosolic
1894	15663	NM_031318	General	testis expressed 1 (Tctex1), mRNA. Length = 698 Rattus norvegicus prolyl	t-complex testis expressed 1
1895	1422	NM_031324	bb, General	endopeptidase (Prep), mRNA. Length = 2743 Rattus norvegicus UDP-	prolyl endopeptidase
1896	18597	NM_031325	g,bb	glucose dehydrogeanse (Ugdh), mRNA. Length = 2318	UDP-glucose dehydrogeanse
1897	11259	NM_031327	i,cc, General	Rattus norvegicus cysteine rich protein 61 (Cyr61), mRNA. Length = 1871	cysteine rich protein 61
				Rattus norvegicus heterogeneous nuclear ribonucleoprotein A/B (Hnrpab), mRNA. Length =	heterogeneous nuclear
1898	4235	NM_031330	General	3061 Rattus norvegicus proteasome (prosome, macropain) 26S subunit, non-	ribonucleoprotein A/B
1899	18375	NM_031331	l,m	ATPase,4 (Psmd4), mRNA. Length = 1334 Rattus norvegicus E-	proteasome (prosome, macropain) 26S subunit, non-ATPase,4
1900	3519	NM_031334	СС	cadherin (Cdh1), mRNA. Length = 4396	E-cadherin
1901	20698	NM 031357	b	Rattus norvegicus ceroid- lipofuscinosis, neuronal 2 (Cln2), mRNA. Length = 2485	
	634			Rattus norvegicus Glutathione-S-transferase, alpha type (Ya) (Gsta1), mRNA. Length = 1178	Glutathione-S-transferase, alpha type
1903	034	NM_031509	n	Rattus norvegicus Glutathione-S-transferase, alpha type (Ya) (Gsta1),	(Ya) Glutathione-S-transferase, alpha type
1903	25525	NM_031509	n	mRNA. Length = 1178 Rattus norvegicus	(Ya)
1903	25069	NM_031509	b,n,w	Glutathione-S-transferase, alpha type (Ya) (Gsta1), mRNA. Length = 1178	
1003	625	NM 024500	7	Rattus norvegicus Glutathione-S-transferase, alpha type (Ya) (Gsta1),	Glutathione-S-transferase, alpha type
1903	635	NM_031509	Z	mRNA. Length = 1178 Rattus norvegicus Met proto- oncogene (Met), mRNA.	(Ya)
1904	848	NM_031517	t	Length = 4189 Rattus norvegicus Nerve growth factor, gamma	Met proto-oncogene
1905	1872	NM_031523	а	polypeptide (Ngfg), mRNA. Length = 873	Nerve growth factor, gamma polypeptide

Table1: 8	UMMARY				/ Ally. Docket No. 44921-5059WG Doc. No. 1793397.
Sequence ID No.	ldeniller	Cenenk Ace Ref. Seg ID	Model Gode	Gene Name	Unigene Guster Title 🗼 🐇
1905	16245	NM_031523	a,d,u	Rattus norvegicus Nerve growth factor, gamma polypeptide (Ngfg), mRNA. Length = 873 Rattus norvegicus Nerve	Rattus norvegicus (clone RSKG50) kallikrein mRNA, 3' end
1905	16244	NM_031523	a	growth factor, gamma polypeptide (Ngfg), mRNA. Length = 873 Rattus norvegicus Protein	Rattus norvegicus (clone RSKG50) kallikrein mRNA, 3' end
1906	9370	NM 031527	w	phosphatase type 1 alpha, catalytic subunit (Ppp1ca), mRNA. Length = 1392	Protein phosphatase type 1 alpha, catalytic subunit
1907	20448	NM 031530	General	Rattus norvegicus Small inducible gene JE (Scya2), mRNA. Length = 780	Small inducible gene JE
1907	20449	NM 031530	General	Rattus norvegicus Small inducible gene JE (Scya2), mRNA. Length = 780	Small inducible gene JE
1908	14633	NM 031533	u	Rattus norvegicus Androsterone UDP- glucuronosyltransferase (Ugt2b2), mRNA. Length = 1593	Androsterone UDP- glucuronosyltransferase
1909	16048	NM_031541	f	Rattus norvegicus CD36 antigen (collagen type I receptor, thrombospondin receptor)-like 1 (scavanger receptor class B type 1) (Cd36I1), mRNA. Length = 2497	CD36 antigen (collagen type I receptor, thrombospondin receptor)-like 1 (scavanger receptor class B type 1)
1910	4011	NM 031543	c,q	Rattus norvegicus Cytochrome P450, subfamily 2e1 (ethanol-inducible) (Cyp2e1), mRNA. Length = 1624	Cytochrome P450, subfamily 2e1 (ethanol-inducible)
1910	4010	NM 031543	c,q	Rattus norvegicus Cytochrome P450, subfamily 2e1 (ethanol-inducible) (Cyp2e1), mRNA. Length = 1624	Cytochrome P450, subfamily 2e1
1910	4012	NM 031543	q	Rattus norvegicus Cytochrome P450, subfamily 2e1 (ethanol-inducible) (Cyp2e1), mRNA. Length = 1624	Cytochrome P450, subfamily 2e1 (ethanol-inducible)
1911	28	NM_031546	General	Rattus norvegicus Regucalcin (Rgn), mRNA. Length = 1605	Regucalcin
1912	24640	NM_031548	h,cc	Rattus norvegicus Sodium channel, nonvoltage-gated 1, alpha (epithelial) (Scnn1a), mRNA. Length = 3081	Sodium channel, nonvoltage-gated 1, alpha (epithelial)
1913	17149	NM_031549	x	Rattus norvegicus Transgelin (Smooth muscle 22 protein) (Tagln), mRNA. Length = 1186	Transgelin (Smooth muscle 22 protein)
			7	Rattus norvegicus Transgelin (Smooth muscle 22 protein) (Tagln), mRNA. Length =	Transgelin (Smooth muscle 22
1913	13105	NM_031549 NM_031552	w	1186 Rattus norvegicus Adducin 3, gamma (Add3), mRNA. Length = 2246	protein) Adducin 3, gamma

TABLE 0: .					AMY, Docker No. 44921-5039W0 Dock No. 1793397.1
Sequence ID No:	(Identifier	GenBenk (Acc) Ref. Seq.(D	Model Gode	Ceno Namo	Unigene Cluster Title
1915	15411	NM_031559	d,r	Length = 4377	Carnitine palmitoyltransferase 1 alpha, liver isoform
1916	16164	NM_031563	а,у	Rattus norvegicus Y box protein 1 (Ybx1), mRNA. Length = 1489	Y box protein 1
1917	9621	NM_031570	bb	Rattus norvegicus ribosomal protein S7 (Rps7), mRNA. Length = 650	ribosomal protein S7
1917	9620	NM_031570	w,bb	Rattus norvegicus ribosomal protein S7 (Rps7), mRNA. Length = 650	ribosomal protein S7
1918	546	NM_031573	f	Rattus norvegicus Phosphorylase kinase, gamma 1 (Phkg1), mRNA. Length = 1388	phosphorylase kinase gamma
1919	1921	NM 031576	f	Rattus norvegicus P450 (cytochrome) oxidoreductase (Por), mRNA. Length = 2441	
				Rattus norvegicus P450 (cytochrome) oxidoreductase	
1919	1920 24219	NM_031576	i.General	Rattus norvegicus protein tyrosine phosphatase 4a1 (Ptp4a1), mRNA. Length = 2638	P450 (cytochrome) oxidoreductase protein tyrosine phosphatase 4a1
1921	770	NM_031584	k,x	Rattus norvegicus solute carrier family 22, member 2 (Slc22a2), mRNA. Length = 2152	solute carrier family 22, member 2
1922	18008	NM_031588	сс	Rattus norvegicus neuregulin 1 (Nrg1), mRNA. Length = 3272	potassium channel, subfamily K, member 3
1922	18005	NM_031588	h	Rattus norvegicus neuregulin 1 (Nrg1), mRNA. Length = 3272	potassium channel, subfamily K, member 3
1922	18011	NM_031588	cc, General	Rattus norvegicus neuregulin 1 (Nrg1), mRNA. Length = 3272	potassium channel, subfamily K, member 3
4000	4504	NIM 024505	L	Rattus norvegicus proteasome (prosome, macropain) 26S subunit, ATPase 3 (Psmc3), mRNA.	proteasome (prosome, macropain) 26S subunit. ATPase 3
1923	1584 24235	NM_031595	k	Length = 1627 Rattus norvegicus thioredoxin reductase 1 (Txnrd1), mRNA. Length = 3360	thioredoxin reductase 1
1924	24234	NM_031614	General	Rattus norvegicus thioredoxin reductase 1 (Txnrd1), mRNA. Length = 3360	thioredoxin reductase 1
				Rattus norvegicus nuclear receptor subfamily 1, group H, member 3 (Nr1h3),	nuclear receptor subfamily 1, group H,
1925	1639	NM_031627	j,l,v m,	mRNA. Length = 1723 Rattus norvegicus core promoter element binding protein (Copeb), mRNA.	member 3
1926	1727	NM_031642	General	Length = 1356	core promoter element binding protein

TABLE 118 S	UMMARY				Ally, Dooket No. 44921-5039W0 Dos. No. 1793397.1
Sequence (D No. : : :	legantijar	ConBank Acc/ Ref. Seg ID	Model Code	Cene Namo	Univers Civeter Tille
		Approximation of the control of the	and the state of the state of	Rattus norvegicus mitogen activated protein kinase kinase 2 (Map2k2), mRNA.	mitogen activated protein kinase
1927	20766	NM_031643	y k,l,m,	Length = 1182 Rattus norvegicus latexin	kinase 2
1929	1993	NM_031655	General	Rattus norvegicus cyclic	latexin
1930	2057	NIM 024660	e	AMP phosphoprotein, 19kD (Arpp19-pending), mRNA. Length = 339	cyclic AMP phosphoprotein, 19kD
1950	2057	NM_031660		Rattus norvegicus solute carrier family 15 (H+/peptide transporter), member 2	
1931	15039	NM_031672	k,General	(Slc15a2), mRNA. Length = 3923	solute carrier family 15 (H+/peptide transporter), member 2
1932	15175	NM 031682	bb	Rattus norvegicus hydroxyacyl-Coenzyme A dehydrogenase, type II (Hadh2), mRNA. Length = 917	hydroxyacyl-Coenzyme A dehydrogenase, type II
1933	1004	NM_031685	v	Rattus norvegicus golgi SNAP receptor complex member 2 (Gosr2), mRNA. Length = 683	golgi SNAP receptor complex member
1934	19727	NM_031687	a,q,s	Rattus norvegicus ubiquitin A 52 residue ribosomal protein fusion product 1 (Uba52), mRNA. Length = 467	ubiquitin A-52 residue ribosomal protein fusion product 1
1935	20404	NM_031700	j.r.y	Rattus norvegicus claudin 3 (Cldn3), mRNA. Length = 1192	claudin 3
1935	20405	NM_031700	o,r	Rattus norvegicus claudin 3 (Cldn3), mRNA. Length = 1192	claudin 3
1936	811	NM_031705	General	Rattus norvegicus dihydropyrimidinase (Dpys), mRNA. Length = 2091	dihydropyrimidinase
1936	812	NM_031705	o,v,bb, General	Rattus norvegicus dihydropyrimidinase (Dpys), mRNA. Length = 2091	dihydropyrimidinase
1937	16204	NM_031706	q,bb	Rattus norvegicus ribosomal protein S8 (Rps8), mRNA. Length = 696	ribosomal protein S8
1937	16205	NM_031706	а,у	Rattus norvegicus ribosomal protein S8 (Rps8), mRNA. Length = 696	ribosomal protein S8
				Rattus norvegicus glycoprotein 110 (Gp110- pending), mRNA. Length =	
1938	24081	NM_031708	m	Rattus norvegicus ribosomal	glycoprotein 110
1939	16918	NM_031709	a,q	protein S12 (Rps12), mRNA. Length = 499	ribosomal protein S12
1940	1081	NM_031712	General	Rattus norvegicus PDZ domain containing 1 (Pdzk1), mRNA. Length = 2005	PDZ domain containing 1
			b,n,u,cc,	Rattus norvegicus phosphofructokinase, muscle (Pfkm), mRNA. Length =	
1941	1340	NM_031715	General	2757 Rattus norvegicus alcohol dehydrogenase family 3, subfamily A2 (Aldh3a2),	phosphofructokinase, muscle alcohol dehydrogenase family 3,
1942	23884	NM_031731	j,s	mRNA. Length = 2977	subfamily A2

TABLE 18 8	DUMMARY				Ally. Dockel No. 44921-5139W0 Doc. No. 1798897.1
Séquince ID'No.	ldeniller	Confinit Acc	Modal : Code	Cono Namo	Unitem Cluster Title
Tomat or what present street any action				Rattus norvegicus UDP- Gal:betaGlcNAc beta 1,4- galactosyltransferase, polypeptide 6 (B4galt6),	UDP-Gal:betaGlcNAc beta 1,4-
1943	10241	NM_031740	d	mRNA. Length = 5729 Rattus norvegicus solute	galactosyltransferase, polypeptide 6
1044	1214	NIM 021741		carrier family 2 (facilitated glucose transporter), member 5 (Slc2a5), mRNA.	solute carrier family 2 (facilitated glucose transporter), member 5
1944	1214	NM_031741	r	Length = 2169 Rattus norvegicus solute carrier family 2 (facilitated glucose transporter),	
1944	1215	NM_031741	r	member 5 (Slc2a5), mRNA. Length = 2169	solute carrier family 2 (facilitated glucose transporter), member 5
1945	20724	NM_031753	h	Rattus norvegicus activated leukocyte cell adhesion molecule (Alcam), mRNA. Length = 2866	activated leukocyte cell adhesion molecule
1946	20753	NM 031763	h	Rattus norvegicus platelet- activating factor acetylhydrolase beta subunit (PAF-AH beta) (Pafah1b1), mRNA. Length = 1233	platelet-activating factor acetylhydrolase beta subunit (PAF-AH beta)
1940	20733	1NW 031703		Rattus norvegicus platelet- activating factor acetylhydrolase beta subunit	platelet-activating factor
1946	20752	NM_031763	у	(PAF-AH beta) (Pafah1b1), mRNA. Length = 1233 Rattus norvegicus rab	acetylhydrolase beta subunit (PAF-AH beta)
				acceptor 1 (prenylated) (Rabac1), mRNA. Length =	
1947	14953	NM_031774	р	Rattus norvegicus guanine	rab acceptor 1 (prenylated)
1948	14184	NM_031776	t,General	deaminase (Gda), mRNA. Length = 1568 Rattus norvegicus guanine	guanine deaminase
4040	14405	NN4 02477C	d,o,t,	deaminase (Gda), mRNA.	
1948	14185	NM_031776	General	Rattus norvegicus NF-E2-	guanine deaminase
1949	1169	NM_031789	С	related factor 2 (Nfe2l2), mRNA. Length = 2307	NF-E2-related factor 2
				Rattus norvegicus defensin beta 1 (Defb1), mRNA.	
1950	16155	NM_031810	d,z	Rattus norvegicus defensin	defensin beta 1
1950	16156	NM_031810	d	beta 1 (Defb1), mRNA. Length = 416	defensin beta 1
				Rattus norvegicus G protein- coupled receptor kinase- associated ADP ribosylation factor GTPase-activating	G protein-coupled receptor kinase-
1951	17194	NM_031814	z	Length = 3236	associated ADP ribosylation factor GTPase-activating protein (GIT1)
				Rattus norvegicus retinoblastoma binding protein 7 (Rbbp7), mRNA.	
1952	17535	NM_031816	bb	Length = 1947 Rattus norvegicus serum-	retinoblastoma binding protein 7
1953	2655	NM_031821	i,l,m,aa	inducible kinase (Snk), mRNA. Length = 2781	serum-inducible kinase
1954	10167	NM_031830	i	Rattus norvegicus reggie1-1 (Flot2), mRNA. Length = 2629	reggie1-1

VABLE 1:	SUMMARY			The state of the s	Aiiy. Dockei No. 44921-5089W0 Doc. No. 1798897.1
Sequate: ID No.	ldenWiter	Genbank Ace/ Ref. Seq [D.	Model Gode	Cons Namo	Unigare Cluster IIIVle
1955	22321	NM_031832	o,t,u, General	Rattus norvegicus IgE binding protein (Lgals3), mRNA. Length = 948 Rattus norvegicus	IgE binding protein
				sulfotransferase family 1A, phenol-preferring, member 1 (Sult1a1), mRNA. Length =	
1956	4748	NM_031834	e,t_	Rattus norvegicus sulfotransferase family 1A, phenol-preferring, member 1	minoxidil sulfotransferase
1956	4749	NM_031834	e,t	(Sult1a1), mRNA. Length = 1227 Rattus norvegicus beta-	minoxidil sulfotransferase
1957	7914	NM_031835	е	alanine-pyruvate aminotransferase (AGT2), mRNA. Length = 2151 Rattus norvegicus vascular	beta-alanine-pyruvate aminotransferase
1958	8385	NM_031836	h	endothelial growth factor (Vegf), mRNA. Length = 645 Rattus norvegicus vascular	vascular endothelial growth factor
1958	8384	NM_031836	h	endothelial growth factor (Vegf), mRNA. Length = 645 Rattus norvegicus ribosomal	vascular endothelial growth factor
1959	10268	NM_031838	а	protein S2 (Rps2), mRNA. Length = 819 Rattus norvegicus ribosomal	ribosomal protein S2
1959	10269	NM_031838	aa	protein S2 (Rps2), mRNA. Length = 819 Rattus norvegicus ribosomal	ribosomal protein S2
1959	10267	NM_031838	n,aa	protein S2 (Rps2), mRNA. Length = 819 Rattus norvegicus stearoyl-	ribosomal protein S2
1960	15077	NM_031841	b	CoA desaturase 2 (Scd2), mRNA. Length = 5055 Rattus norvegicus	stearoyl-CoA desaturase 2
1961	16726	NM_031855	x	Ketohexokinase (Khk), mRNA. Length = 1342	Ketohexokinase
1962	25802	NM_031969	а	Rattus norvegicus Calmodulin 1 (phosphorylase kinase, delta) (Calm1), mRNA. Length = 3513	Calmodulin 1 (phosphorylase kinase, delta)
				Rattus norvegicus Calmodulin 1 (phosphorylase kinase, delta) (Calm1),	Calmodulin 1 (phosphorylase kinase,
1962	19191	NM_031969	С	mRNA. Length = 3513 Rattus norvegicus Calmodulin 1 (phosphorylase	delta)
1962	19195	NM_031969	r	kinase, delta) (Calm1), mRNA. Length = 3513	Calmodulin 1 (phosphorylase kinase, delta)
1962	19190	NM_031969	p	Rattus norvegicus Calmodulin 1 (phosphorylase kinase, delta) (Calm1), mRNA. Length = 3513	Calmodulin 1 (phosphorylase kinase, delta)
1963	17734	NM_031970	v,General	Rattus norvegicus Heat shock 27 kDa protein (Hsp27), mRNA. Length = 787	ESTs,Heat shock 27 kDa protein
1964	1475	NM 031971	v,Gerierai	Rattus norvegicus Heat	ESTs, Highly similar to S10A RAT S- 100 PROTEIN, ALPHA CHAIN [R.norvegicus], Heat shock protein 70- 1

TABUEN: 8) Ally, Docket No. 44921-5039W@ Doc. No. 1798697.1
Sequence ID No.	ldentifier	Genbank Acci Rei. Seg ID	Model Code	Cene Name	Unigane Gluster Title
1965	15470	NM_031978	f	Rattus norvegicus 26S proteasome, subunit p112 (PSMD1), mRNA. Length = 3089	26S proteasome, subunit p112
1966	18502	NM 031984	С	Rattus norvegicus cerebellar Ca-binding protein, spot 35 protein (Calb1), mRNA. Length = 2280	cerebellar Ca-binding protein, spot 35 protein
1967	19768	NM_031986	v,aa, General	Rattus norvegicus syntenin (Sdcbp), mRNA. Length = 2077	syntenin
1968	723	NM_032084	n	Rattus norvegicus chimerin (chimaerin) 2 (Chn2), mRNA. Length = 1118 Rattus norvegicus	chimerin (chimaerin) 2
1969	17935	NM_032615	a	membrane interacting protein of RGS16 (Mir16), mRNA. Length = 1203	membrane interacting protein of RGS16
1970	16831	NM 033095	n	Rattus norvegicus Crystallin, gamma polypeptide 4 (Crygd), mRNA. Length = 634	
1971	25468	NM_033234	c,z	Rattus norvegicus Hemoglobin, beta (Hbb), mRNA. Length = 620	
1971	25469	NM_033234	С	Rattus norvegicus Hemoglobin, beta (Hbb), mRNA. Length = 620 Rattus norvegicus	
1971	17832	NM_033234	с,р	Hemoglobin, beta (Hbb), mRNA. Length = 620 Rattus norvegicus	Rat major beta-globin mRNA, complete cds
1971	17829	NM_033234	c,z	Hemoglobin, beta (Hbb), mRNA. Length = 620 Rattus norvegicus Malate	Rat major beta-globin mRNA, complete cds
1972	4723	NM_033235	z	dehydrogenase-like enzyme (Mdhl), mRNA. Length = 1266 Rattus norvegicus	Rattus norvegicus cytosolic malate dehydrogenase (Mdh) mRNA, complete cds
1973	1409	NM_033349	p,General	Hydroxyacyl glutathione hydrolase (Hagh), mRNA. Length = 783	Rattus norvegicus round spermatid protein RSP29 gene, complete cds
				Rattus norvegicus ATP- binding cassette, sub-family D (ALD), member 2 (Abcd2),	
1974 1975	19998	NM_033352	General d	mRNA. Length = 5531 Rattus norvegicus Kidney 1 (Kid1), mRNA. Length = 2563	PDZ domain containing 1 Rat zinc finger protein (kid-1) mRNA, complete cds
1373	1410	NM_052798	-	Rattus norvegicus cytosolic cysteine dioxygenase 1 (Cdo1), mRNA. Length =	Rat cysteine dioxygenase mRNA,
1976	15028	NM_052809	f	Rattus norvegicus Pyruvate kinase 3 (Pkm2), mRNA.	complete cds Rat mRNA for pituitary pyruvate
1977	7660	NM_053297 NM_053299	i	Length = 1973 Rattus norvegicus ubiquitin D (Ubd), mRNA. Length = 684	kinase ESTs, Weakly similar to polyubiquitin [R.norvegicus]
1979	5117	NM_053310	p_	Rattus norvegicus homer, neuronal immediate early gene, 3 (Homer3), mRNA. Length = 1207	Rattus norvegicus mRNA for Vesl-3, complete cds

Sequence GenPank Ace Modal Gene Modal Gene Modal Gene Modal Gene Modal Gene Modal Gene Modal Gene Modal Gene Modal Gene Modal Gene Modal Gene G	21-5089W0 . 1798997.1
1981 17473	
1982 25480 NM 053329 g g g mRNA. Length = 1812 Rattus norvegicus insulin-like growth factor binding protein, acid labile subunit (gfals), mRNA. Length = 1812 Rattus norvegicus insulin-like growth factor binding protein complete complet	
growth factor binding protein, acid labile subunit (lgfals), mRNA. Length = 1812 Rattus norvegicus insulinding protein complete comple	
1983 14926	plex acid-
1983 14929 NM_053330 e,General Langh = 554 Eattus norvegicus cribosom L21 mRNA, complete cds Rattus norvegicus cubilin (intrinsic factor-cobalamin receptor) (Cubn), mRNA. Rattus norvegicus intrinsic receptor) (Cubn), mRNA. Rattus norvegicus ribosom L21 mRNA, complete cds Rattus norvegicus cubilin (intrinsic factor-cobalamin receptor) (Cubn), mRNA. Rattus norvegicus intrinsic receptor) (Cubn), mRNA. Length = 10.872 Complete cds Rattus norvegicus regulator of G-protein signaling 19 (Rgs 19), mRNA. Length = 1607 Rattus norvegicus interferon gamma inducing factor binding protein (lgiftp), mRNA. Length = 2 (Col1a2), mRNA. Length = 626 Rattus norvegicus solute carrier family 34 (sodium phosphate), member 2 (Slc34a2), mRNA. Length = 2 (Slc34a2), mRNA. Length = 1988 6416 NM_053380 General 3950 Rattus norvegicus small muscle protein, X-linked (Smpx), mRNA. Length = 1989 19113 NM_053395 Rattus norvegicus flavincontaining monooxygenase 3 (Fmo3), mRNA. Length = 1990 2242 NM_053433 n,General 2037 Rattus norvegicus zinc finger protein 103 (Zfp103), mRNA. Length = 1047 Rattus norvegicus zinc finger protein 103 (Zfp103), mRNA. Length = 1047 Rattus norvegicus zinc finger protein 103 (Zfp103), mRNA. Length = 1047 Rattus norvegicus zinc finger protein 103 (Zfp103), mRNA. Length = 1047 Rattus norvegicus zinc finger protein 103 (Zfp103), mRNA. Length = 1047 Rattus norvegicus zinc finger protein 103 (Zfp103), mRNA. Length = 1047 Rattus norvegicus zinc finger protein 103 (Zfp103), mRNA. Length = 1047 Rattus norvegicus zinc finger protein 103 (Zfp103), mRNA. Length = 1047 Rattus norvegicus zinc finger protein 103 (Zfp103), mRNA. Length = 1047 Rattus norvegicus zinc finger protein 103 (Zfp103), mRNA. Length = 1047 Rattus norvegicus zinc finger protein 103 (Zfp103), mRNA. Length = 1047 Rattus norvegicus zinc finger protein 103 (Zfp103), mRNA. Length = 1047 Rattus norvegicus zinc finger protein 103 (Zfp103), mRNA. Length = 1047 Rattus norvegicus zin	
(intrinsic factor-cobalamin receptor) (Cubn), mRNA. In the process of G-protein signaling 19 (Rgs19), mRNA. Length = 10,872 (Collagen, type I, alpha 2 (Collagen, type I, alpha 2 (Collagen, type I, alpha 2 (Collagen, type I), a	
1985 15790 NM 053341 j,x 1607 regulator of G-protein signaling 19 (Rgs19), mRNA. Length = 1607 regulator of G-protein signaling 19 (Rgs19), mRNA. Length = 1607 regulator of G-protein signaling 19 (Rgs19), mRNA. Length = 1607 regulator of G-protein signaling 19 (Col1a2), mRNA. Length = 1607 regulator of G-protein signaling 19 (Col1a2), mRNA. Length = 1607 regulator of G-protein signaling 19 (Rattus norvegicus interferon gamma inducing factor binding protein (Igifbp), interferon gamma inducing formal inducing factor binding protein (Igifbp), interferon gamma inducing protein (Igifbp), interferon gamma inducing binding protein interferon gamma inducing binding protein interferon gamma inducing binding protein interferon gamma inducing binding protein interferon gamma inducing interferon gamma inducing binding protein interferon gamma inducing binding protein interferon gamma inducing interferon gamma inducing binding protein interferon gamma inducing binding protein interferon gamma inducing interferon gamma inducing binding protein interferon gamma inducing interferon gamma inducing side interferon gamma inducing interferon gamma inducing binding protein interferon gamma inducing interferon gamma indu	
Rattus norvegicus procollagen, type I, alpha 2 (Cof1a2), mRNA. Length = 4474 Procollagen, type I, alpha 2 (Cof1a2), mRNA. Length = procollagen, type I, alpha 2 (Cof1a2), mRNA. Length = procollagen, type I, alpha 2 (Cof1a2), mRNA. Length = procollagen, type I, alpha 2 (Rattus norvegicus interferon gamma inducing factor binding protein (Igifbp), interferon gamma inducing protein (Igifbp), mRNA. Length = 626 (Sic34a2), mRNA. Length = solute carrier family 34 (sodium phosphate), member 2 (Sic34a2), mRNA. Length = solute carrier family 34 (so phosphate), member 2 (Sic34a2), mRNA. Length = Rattus norvegicus small muscle protein, X-linked (Smpx), mRNA. Length = Rattus norvegicus SMPX procontaining monooxygenase 3 (Fmo3), mRNA. Length = flavin-containing monooxygenase 3 (Fmo3), mRNA. Leng	
1986 6154 NM_053356 p 4474 procollagen, type I, alpha 2 Rattus norvegicus interferon gamma inducing factor binding protein (Igifbp), interferon gamma inducing binding protein (Igifbp), mRNA. Length = 626 Rattus norvegicus solute carrier family 34 (sodium phosphate), member 2 (SIc34a2), mRNA. Length = 3950 Rattus norvegicus small muscle protein, X-linked (Smpx), mRNA. Length = Rattus norvegicus SMPX p. (Smpx) mRNA. Length = Rattus norvegicus SMPX p. (Smpx) mRNA. Length = 1990 Rattus norvegicus flavincontaining monooxygenase 3 (Fmo3), mRNA. Length = 1990 Rattus norvegicus zinc finger protein 103 (Zfp103), mRNA. Length = 100 (Zfp103), mRNA. Length =	ialing 19
binding protein (Igifbp), mRNA. Length = 626 binding protein Rattus norvegicus solute carrier family 34 (sodium phosphate), member 2 (Slc34a2), mRNA. Length = 3950 solute carrier family 34 (sodium phosphate), member 2 Rattus norvegicus small muscle protein, X-linked (Smpx), mRNA. Length = 892 (Smpx) mRNA. Length = 892 (Smpx) mRNA. Length = 892 (Smpx) mRNA. Complete of Smpx) mRNA. Length = 892 (Smpx) mRN	2
Carrier family 34 (sodium phosphate), member 2 (Slc34a2), mRNA. Length = 3950 Solute carrier family 34 (so phosphate), member 2 (Slc34a2), mRNA. Length = 3950 Solute carrier family 34 (so phosphate), member 2 Solute carrier family 34 (solute family 34 (solute family 34 (solute family 34 (solute family 34 (solute family 3	g factor
muscle protein, X-linked (Smpx), mRNA. Length = Rattus norvegicus SMPX protein (Smpx) mRNA. Length = Rattus norvegicus SMPX protein 103 (Zfp103), mRNA. Length = 1990	odium
containing monooxygenase 3 (Fmo3), mRNA. Length = 1990 2242 NM_053433 n,General 2037 flavin-containing monooxygenase 2037 flavin-containing monooxygenase 7 (Factor of the containing	
Rattus norvegicus zinc finger protein 103 (Zfp103), mRNA. 1991 5561 NM_053438 y Length = 3258 zinc finger protein 103	
1991 5561 NM_053438 y Length = 3258 zinc finger protein 103	genase 3
member RAS oncogene	
1992 14670 NM_053439 N,General = 1084 RAN, member RAS oncoger Rattus norvegicus	ene family
superiorcervical ganglia, neural specific 10 (Scgn10), superiorcervical ganglia, neural specific 10 (Scgn10), specific 10 1993 17102 NM_053440 w mRNA. Length = 1654 specific 10	ieural
Rattus norvegicus solute carrier family 8 (cationic amino acid transporter, y+ system), member 7 (Lat4), acid transporter, y+ system 1994 24762 NM 053442 General mRNA. Length = 4117 7	

TABLET: S	WMMARY				ANN. Docket No. 44921-5039WO Doc, No. 1793397.1
Sequence ID No.	ldentifier	Consonk Aco Ref. Seq ID	(Model))	Cene Namo	Unigene Cluster Tillo
1995	8085	NM_053453	General	Rattus norvegicus regulator of G-protein signaling protein 2 (Rgs2), mRNA. Length = 1629	regulator of G-protein signaling protein 2
1996	4622	NM_053463	d	Rattus norvegicus nucleobindin (Nucb), mRNA. Length = 2303	nucleobindin
1997	21866	NM 053472	p	Rattus norvegicus cytochrome c oxidase subunit IV isoform 2 precursor (CoxIV-2), mRNA. Length = 704	cytochrome c oxidase subunit IV isoform 2 precursor
1998	9573	NM 053475	h	Rattus norvegicus protein tyrosine phosphatase type IVA, member 2 (Ptp4a2), mRNA. Length = 1095	protein tyrosine phosphatase type IVA, member 2
1999	16137	NM_053480	k	Rattus norvegicus DNA polymerase alpha subunit II (Pola2), mRNA. Length = 1836	DNA polymerase alpha subunit II
2000	15556	NM_053483	у	Rattus norvegicus karyopherin (importin) alpha 2 (Kpna2), mRNA. Length = 1886 Rattus norvegicus calcium	karyopherin (importin) alpha 2
2001	16394	NM_053485	General	binding protein A6 (calcyclin) (S100a6), mRNA. Length = 291 Rattus norvegicus	calcium binding protein A6 (calcyclin)
2002	4290	NM 053487	j,y	peroxisomal membrane protein Pmp26p (Peroxin-11) (Pex11a), mRNA. Length = 1194	peroxisomal membrane protein Pmp26p (Peroxin-11)
2004	18826	NM 053523	d	Rattus norvegicus homocysteine-inducible, endoplasmic reticulum stress inducible, ubiquitin-like domain member 1 (Herpud1), mRNA. Length = 1857	
2005	7764	NM 053525	aa	Rattus norvegicus ATP- dependent, RNA helicase (Rok1), mRNA. Length = 2175	Rattus norvegicus rROK1L mRNA for ROK1-like protein, complete cds
				Rattus norvegicus lysosomal- associated protein transmembrane 5 (Laptm5),	Rattus norvegicus gcd-10S mRNA,
2006	14199	NM_053538	С	mRNA. Length = 1309 Rattus norvegicus isopentenyl-diphosphate delta isomerase (Idi1),	Complete cds Rattus norvegicus isopentenyl diphosphate:dimethylallyl diphosphate
2007	1058	NM_053539 NM_053563	c,d General	mRNA. Length = 1182 Rattus norvegicus nuclear RNA helicase, DECD variant of DEAD box family (Ddxl), mRNA. Length = 1511	Rattus norvegicus nuclear RNA helicase mRNA, complete cds
2009	1342	NM_053573	h	Rattus norvegicus olfactomedin related ER localized protein (Olfm1), mRNA. Length = 2759	Rattus norvegicus neuronal olfactomedin-related ER localized protein (D2Sut1e) mRNA, complete cds
2010	19254	NM_053576	h,s	Rattus norvegicus thiol- specific antioxidant protein (Prdx5), mRNA. Length = 1414	Rattus norvegicus mRNA for thiol- specific antioxidant protein (1-Cys peroxiredoxin)

PCT/US02/16173

TIABLE 18 S	UMMARY	4. 渊源			ÁMy, Dockel No. 44921-5039WC Doc, No. 1793897.1
Seguence Seguence	Mentiler	ConBenk Ace Rol Sog ID	Model Gode	Cono Namo	Unigene Civeter Tille
2010	19253	NM_053576	h	Rattus norvegicus thiol- specific antioxidant protein (Prdx5), mRNA. Length = 1414	Rattus norvegicus mRNA for thiol- specific antioxidant protein (1-Cys peroxiredoxin)
2011	3049	NM 053582	p,cc, General	Rattus norvegicus glucocorticoid-inducible protein (gis5), mRNA. Length = 1869	Rattus norvegicus gis5 mRNA for glucocorticoid-inducible protein, complete cds
			o General	Rattus norvegicus glucocorticoid-inducible protein (gis5), mRNA. Length = 1869	Rattus norvegicus gis5 mRNA for glucocorticoid-inducible protein, complete cds
2011	21423	NM_053582 NM_053586	o,General	Rattus norvegicus cytochrome c oxidase subunit Vb (Cox5b), mRNA. Length = 485	Rat mRNA for cytochrome c oxidase subunit VIa
2013	21445	NM_053587	t,v	Rattus norvegicus S100 calcium-binding protein A9 (calgranulin B) (S100a9), mRNA. Length = 494	Rattus norvegicus intracellular calcium binding protein (MRP14) mRNA, complete cds
2014	20871	NM_053591	j,l	Rattus norvegicus dipeptidase 1 (Dpep1), mRNA. Length = 2179 Rattus norvegicus	Rat dipeptidase (dpep1) mRNA, complete cds
2014	20870	NM_053591	1	dipeptidase 1 (Dpep1), mRNA. Length = 2179 Rattus norvegicus protein	Rat dipeptidase (dpep1) mRNA, complete cds
2015	21044	NM_053594	d	tyrosine phosphatase, receptor type, R (Ptprr), mRNA. Length = 3565	Rattus norvegicus mRNA for tyrosine phosphatase CBPTP, complete cds
2016	21709	NM_053596	k	Rattus norvegicus endothelin- converting enzyme 1 (Ece1), mRNA. Length = 4469	Rat mRNA for endothelin-converting enzyme, complete cds
2016	21708	NM_053596	z	Rattus norvegicus endothelin- converting enzyme 1 (Ece1), mRNA. Length = 4469	Rat mRNA for endothelin-converting enzyme, complete cds
2017	1597	NM_053611	t	Rattus norvegicus nuclear proten 1 (Nupr1), mRNA. Length = 602	Rattus norvegicus p8 mRNA, complete cds
2018	5565	NM_053618	General	Rattus norvegicus Bardet- Biedl syndrome 2 (human) (Bbs2), mRNA. Length = 2573	Rattus norvegicus BBS2 (Bbs2) mRNA, complete cds
2019	13004	NM_053623	t	Rattus norvegicus fatty acid- Coenzyme A ligase, long chain 4 (Facl4), mRNA. Length = 4862	Rattus norvegicus mRNA for Acyl- CoA synthetase, complete cds
2020	1127	NM_053626	g	Rattus norvegicus D-amino acid oxidase (Dao1), mRNA. Length = 1646	Rattus norvegicus mRNA for D-amino- acid oxidase, complete cds
2021	18644	NM 053648	n	Rattus norvegicus beta- carotene 15, 15'- dioxygenase (Bcdo), mRNA. Length = 2207	Rattus norvegicus mRNA for beta- carotene 15,15'-dioxygenase, complete cds
				Rattus norvegicus vascular endothelial growth factor C (Vegfc), mRNA. Length =	ESTs, Highly similar to VEGC MOUSE VASCULAR ENDOTHELIAL GROWTH FACTOR C PRECURSOR
2022	3454	NM_053653 NM_053662	СС	1596 Rattus norvegicus cyclin L (Ccnl), mRNA. Length = 2092	[M.musculus] Rattus norvegicus cyclin ania-6a mRNA, complete cds

TABLE 1:	SUMMARY				Atty, Docket No. 44921-5039W0 Doc. No. 1793397.1
Sequence ID No.	ldeniller	GenBank Ace/ Ref. Seq.ID	Model Code	Cene Name	Unijem Gluster Tilie
Total Section			According to the second	Rattus norvegicus Cbp/p300- interacting transactivator, with Glu/Asp-rich carboxy- terminal domain, 2 (Cited2),	
2024	16121	NM_053698	h,j,z	mRNA. Length = 1155 Rattus norvegicus Cbp/p300- interacting transactivator,	MRG1 mRNA, complete cds
2024	16122	NM_053698	h,j,z	with Glu/Asp-rich carboxy- terminal domain, 2 (Cited2), mRNA. Length = 1155 Rattus norvegicus Kruppel-	Rattus norvegicus transcription factor MRG1 mRNA, complete cds
2025	25379	NM_053713	General	like factor 4 (gut) (Klf4), mRNA. Length = 2393 Rattus norvegicus Kruppel-	
2025	13622	NM_053713	General	like factor 4 (gut) (Klf4), mRNA. Length = 2393 Rattus norvegicus ubiquilin 1	ESTs, Moderately similar to zinc finger protein [R.norvegicus]
2026	15376	NM_053747	h	(Ubqln1), mRNA. Length = 2131 Rattus norvegicus dipeptidylpeptidase III	Rattus norvegicus mRNA for DA41, complete cds
2027	1218	NM_053748	b	(Dpp3), mRNA. Length = 2632 Rattus norvegicus	Rattus norvegicus mRNA for dipeptidyl peptidase III, complete cds
2028	1137	NM 053763	V	cytochrome P450, 40 (25- hydroxyvitamin D3 1 alpha- hydroxylase) (Cyp40), mRNA. Length = 2426	Rattus norvegicus 25-hydroxyvitamin D 1-hydroxylase (CYP1) mRNA, complete cds
2028	1137	NW_055765	<u> </u>	Rattus norvegicus protein tyrosine phosphatase, non-receptor type 16 (Ptpn16),	Rattus norvegicus protein tyrosine
2029	15996 8652	NM_053769 NM_053774	g	mRNA. Length = 1104 Rattus norvegicus ubiquitin specific protease 2 (Usp2), mRNA. Length = 1857	phosphatase mRNA, complete cds Rattus norvegicus deubiquitinating enzyme Ubp69 (ubp69) mRNA, complete cds
				Rattus norvegicus potassium channel, subfamily K, member 6 (TWIK-2) (Kcnk6),	
2031	14664 4361	NM_053806 NM_053812	General k	mRNA. Length = 2243 Rattus norvegicus BCL2- antagonist/killer 1 (Bak1), mRNA. Length = 1923	Rattus norvegicus BAK protein (Bak) mRNA, complete cds
2032	15002	NM 053819	b,x,bb, General	Rattus norvegicus tissue	Rattus norvegicus tissue inhibitor of metalloproteinase-1 (TIMP1), mRNA, complete cds
2034	15003	NM_053819	b,I,x,bb, General	Rattus norvegicus tissue inhibitor of metalloproteinase 1 (Timp1), mRNA. Length = 740	
2035	16173	NM 053822	t	Rattus norvegicus S100 calcium-binding protein A8 (calgranulin A) (S100a8), mRNA. Length = 361	Rattus norvegicus intercellular calcium binding protein (MRP8) mRNA, complete cds
2036	17154	NM_053835	j,z	Rattus norvegicus clathrin, light polypeptide (Lcb) (Cltb), mRNA. Length = 982	Rat clathrin light chain (LCB2) mRNA, complete cds,Rat clathrin light chain (LCB3) mRNA, complete cds
2037	20868	NM 053843	t	Rattus norvegicus Fc receptor, IgG, Iow affinity III (Fcgr3), mRNA. Length = 1318	Rat Fc-gamma receptor mRNA, complete cds
2037	20869	NM 053843	t	Rattus norvegicus Fc receptor, IgG, Iow affinity III (Fcgr3), mRNA. Length = 1318	Rat Fc-gamma receptor mRNA, complete cds

TABLE 1: S	UMMARY				Aity: Dockei No. 44921-5089WG Doc. No. 1793897.1
Sequence: ID No.42	ldentiler	Conbent Acel Ref. Seq ID	Model :	Gene Name	Unigeno Cluster Tillo
2040	714	NM 053863	у	Rattus norvegicus solute carrier family 28 (sodium- coupled nucleoside transporter), member 1 (Slc28a1), mRNA. Length = 2401	Rattus norvegicus Sprague-Dawley sodium-dependent nucleoside transporter (rCNT1) mRNA, complete cds
2041	19781	NM 053883	b	Rattus norvegicus dual specificity phosphatase 6 (Dusp6), mRNA. Length = 2104	Rattus norvegicus dual-specificity protein tyrosine phosphatase (rVH6) mRNA, complete cds
2041	19780	NM_053883	b	Rattus norvegicus dual specificity phosphatase 6 (Dusp6), mRNA. Length = 2104	Rattus norvegicus dual-specificity protein tyrosine phosphatase (rVH6) mRNA, complete cds
2042	1454	NM_053887	General	Rattus norvegicus mitogen activated protein kinase kinase 1 (Map3k1), mRNA. Length = 5180	Rattus norvegicus MAP kinase kinase kinase 1 (MEKK1) mRNA, complete cds
2043	1660	NM_053891	g	Rattus norvegicus cyclin- dependent kinase 5, regulatory subunit 1 (p35) (Cdk5r), mRNA. Length = 1208	Rattus norvegicus P35 mRNA, complete cds
2044	712	NM_053896	k	Rattus norvegicus aldehyde dehydrogenase family 1, subfamily A2 (Aldh1a2), mRNA. Length = 2240 Rattus norvegicus	Rattus norvegicus aldehyde dehydrogenase mRNA, complete cds
2045	753	NM_053897	k	coagulation factor II (thrombin) receptor-like 1 (F2rl1), mRNA. Length = 1428	Rattus norvegicus proteinase- activated receptor-2 mRNA, complete cds
2046	794	NM_053902	General	Rattus norvegicus kynureninase (L-kynurenine hydrolase) (Kynu), mRNA. Length = 1765	Rattus norvegicus L-kynurenine hydrolase mRNA, complete cds
2047	17937	NM_053911	f	Rattus norvegicus pleckstrin homology, Sec7 and coiled/coil domains 2 (Pscd2), mRNA. Length = 1561	Rattus norvegicus sec7B mRNA, complete cds
2048	8188	NM 053927	General	Rattus norvegicus erythrocyte membrane protein band 4.1-like 3 (Epb41I3), mRNA. Length = 4543	Rattus norvegicus mRNA for type II brain 4.1, complete cds
				Rattus norvegicus endothelial differentiation, lysophosphatidic acid G- protein-coupled receptor, 2	Rattus norvegicus putative G-protein
2050	1628	NM_053936	h	(Edg2), mRNA. Length = 1543 Rattus norvegicus crystallin,	coupled receptor GPCR91 (Gpcr91) mRNA, complete cds
2051	13954	NM_053955	General	mu (Crym), mRNA. Length = 1227 Rattus norvegicus	Rattus norvegicus CDK108 mRNA
2052	408	NM_053961	General	endoplasmic retuclum protein 29 (Erp29), mRNA. Length = 4529 Rattus norvegicus	R.norvegicus mRNA encoding 45kDa protein which binds to heymann nephritis antigen gp330
2052	19991	NM_053961	а	endoplasmic retuclum protein 29 (Erp29), mRNA. Length = 4529	mitochondrial aconitase (nuclear aco2 gene)

TABLE 1: S	MWWALLA				Ally. Docket No. 44921-5039W0 Doc. No. 1793397.1
ID No. Sectiones	ldenMler	ConBenk Acc	Model Model	Gene Name	Unigene Cluster Tille
2052	16190	NM_053961	q	Rattus norvegicus endoplasmic retuclum protein 29 (Erp29), mRNA. Length = 4529	ESTs, Weakly similar to ECHM RAT ENOYL-COA HYDRATASE, MITOCHONDRIAL PRECURSOR [R.norvegicus]
				Rattus norvegicus endoplasmic retuclum protein 29 (Erp29), mRNA.	
2052	21355	NM_053961	j,l,y,z	Length = 4529 Rattus norvegicus ribosomal protein L6 (Rpl6), mRNA.	ESTs R.norvegicus mRNA for ribosomal
2055	15136_	NM_053971	aa	Length = 963 Rattus norvegicus ribosomal	protein L6
2055	15135	NM_053971	d	protein L6 (Rpl6), mRNA. Length = 963 Rattus norvegicus eukaryotic	R.norvegicus mRNA for ribosomal protein L6
2056	1764	NM_053974	h	translation initiation factor 4E (Eif4e), mRNA. Length = 1647 Rattus norvegicus ADP-	R.norvegicus mRNA eIF-4E
2057	1292	NM_053980	l .	ratius norvegicus ADF- ribosylation factor related protein 1 (Arfrp1), mRNA. Length = 943 Rattus norvegicus ribosomal	R.norvegicus (Sprague Dawley) ARP1 mRNA for ARF-related protein
2058	15468	NM_053982	q	protein S15a (Rps15a), mRNA. Length = 449 Rattus norvegicus H3	R.norvegicus mRNA for ribosomal protein S15a
2059	15642	NM_053985	General	histone, family 3B (H3f3b), mRNA. Length = 1107 Rattus norvegicus CD36	R.norvegicus mRNA for histone H3.3
2060	21066	NM_054001	t	antigen (collagen type I receptor, thrombospondin receptor)-like 2 (Cd36l2), mRNA. Length = 1938	Rat lysosomal membrane protein (LIMPII) mRNA, complete cds
2061	17326	NM_054008	О	Rattus norvegicus Rgc32 protein (Rgc32), mRNA. Length = 889	Rattus norvegicus RGC-32 (RGC-32) mRNA, complete cds
2061	17327	NM_054008	cc	Rattus norvegicus Rgc32 protein (Rgc32), mRNA. Length = 889	Rattus norvegicus RGC-32 (RGC-32) mRNA, complete cds
2061	17329	NM_054008	g,o,cc	Rattus norvegicus Rgc32 protein (Rgc32), mRNA. Length = 889	Rattus norvegicus RGC-32 (RGC-32) mRNA, complete cds
				Rattus norvegicus proteasome (prosome, macropain) subunit, beta type 6 (Psmb6), mRNA.	
2062	25253	NM_057099	j,l,m,p,z	Length = 760 Rattus norvegicus proteasome (prosome, macropain) subunit, beta type 6 (Psmb6), mRNA.	ESTs, Highly similar to PROTEASOME DELTA CHAIN
2062	22849	NM_057099	j,l	Length = 760 Rattus norvegicus A kinase	PRECURSOR [R.norvegicus]
2063	19657	NM_057103	b,cc	(PRKA) anchor protein (gravin) 12 (Akap12), mRNA. Length = 5236 Rattus norvegicus UDP	Rattus norvegicus PKC binding protein and substrate mRNA, complete cds
2064	5492	NM_057105	w	glycosyltransferase 1 family, polypeptide A6 (Ugt1a6), mRNA. Length = 1593	ESTs,UDP-glucuronosyltransferase 1 family, member 1
2064	15126	NM_057105	r	Rattus norvegicus UDP glycosyltransferase 1 family, polypeptide A6 (Ugt1a6), mRNA. Length = 1593	Rattus norvegicus UDP- glucuronosyltransferase UGT1A7 mRNA, complete cds,UDP- glucuronosyltransferase 1 family, member 1

1/4131713-418 8					Ally. Docket No. 44921-5189WG Doc. No. 1793397.1
Sequence In No.	ldenWiter	ConBank Acc/ Ref. Seq [D	Model Code	Cene Name	Unicana Guster Tilla
2064	15125	NM 057105	s	Rattus norvegicus UDP glycosyltransferase 1 family, polypeptide A6 (Ugt1a6), mRNA. Length = 1593	Rattus norvegicus UDP- glucuronosyltransferase UGT1A7 mRNA, complete cds,UDP- glucuronosyltransferase 1 family, member 1
2066	15391	NM_057114	n	Rattus norvegicus peroxiredoxin 1 (Prdx1), mRNA. Length = 882	Rat mRNA for HBP23 (heme-binding protein 23 kDa), complete cds
2067	727	NM 057123	m	Rattus norvegicus protease (prosome, macropain) 26S subunit, ATPase 1 (Psmc1), mRNA. Length = 1556	Rattus norvegicus mRNA for proteasomal ATPase (S4), complete cds
2068	915	NM_057124	s	Rattus norvegicus pyrimidinergic receptor P2Y, G-protein coupled, 6 (P2ry6), mRNA. Length = 1922 Rattus norvegicus	Rat mRNA for novel G protein-coupled P2 receptor, complete cds
2069	15151	NM_057131	k	phosphoribosyl pyrophosphate synthetase- associated protein 2 (Prpsap2), mRNA. Length = 1612	Rattus norvegicus mRNA for 41-kDa phosphoribosylpyrophosphate synthetase-associated protein, complete cds
2070	1892	NM_057144	b	Rattus norvegicus cysteine- rich protein 3 (Csrp3), mRNA. Length = 853	R.norvegicus mRNA for muscle LIM protein
2071	12333	NM 057155	f	Rattus norvegicus X-prolyl aminopeptidase (aminopeptidase P) 2, membrane-bound (Xpnpep2), mRNA. Length = 2828	Rattus norvegicus membrane-bound aminopeptidase P mRNA, complete cds
2071	12331	NM_057155	v,General	Rattus norvegicus X-prolyl aminopeptidase (aminopeptidase P) 2, membrane-bound (Xpnpep2), mRNA. Length =	Rattus norvegicus membrane-bound aminopeptidase P mRNA, complete cds
2071	12332	NM 057155	f,General	Rattus norvegicus X-prolyl aminopeptidase (aminopeptidase P) 2, membrane-bound (Xpnpep2), mRNA. Length = 2828	Rattus norvegicus membrane-bound aminopeptidase P mRNA, complete cds
2072	17477	NM 057194	a,General	Rattus norvegicus phospholipid scramblase 1 (Plscr1), mRNA. Length = 1569	Rattus norvegicus phospholipid scramblase PLSCR mRNA, complete cds
2073	15408	NM_057197	p,t	Rattus norvegicus 2,4- dienoyl CoA reductase 1, mitochondrial (Decr1), mRNA. Length = 1109	Rattus norvegicus mRNA for 2,4- dienoyl-CoA reductase precursor, complete cds
2073	15409	NM_057197	t	Rattus norvegicus 2,4- dienoyl CoA reductase 1, mitochondrial (Decr1), mRNA. Length = 1109	Rattus norvegicus mRNA for 2,4- dienoyl-CoA reductase precursor, complete cds
2074	7866	NM 057198	h	Rattus norvegicus phosphoribosyl pyrophosphate amidotransferase (Ppat), mRNA. Length = 2934	Rattus norvegicus mRNA for amidophosphoribosyltransferase

TABLEN: S	MAINATEA				Atty, Docket No. 44921-5039WC Doc. No. 1793397.1
Sequence ID No.	ldeniller:	Cenbank Acel Ref. Seq ID	Model Gode	Cene Namo	Unigene Cluster VIIIe
2075	14125	NM 057208	h,j,y,z	Rattus norvegicus tropomyosin 3, gamma (Tpm3), mRNA. Length = 1101	Rattus norvegicus tropomyosin non- muscle isoform NM1 (TPM-gamma) mRNA, complete cds,Rattus norvegicus tropomyosin non-muscle isoform NM3 (TPM-gamma) mRNA, complete cds
2076	1743	NM 057210	k,s	Rattus norvegicus synaptic vesicle glycoprotein 2 a (Sv2a), mRNA. Length = 3844	Rattus norvegicus synaptic vesicle protein (SV2) mRNA, complete cds
2077	10498	NM_078617	а	Rattus norvegicus ribosomal protein S23 (Rps23), mRNA. Length = 432	R.norvegicus (Sprague-Dawley) ribosomal protein S23 mRNA
2078	8820	NM_080399	n	Rattus norvegicus Smhs1 protein (Smhs1), mRNA. Length = 1107 Rattus norvegicus ATP-	ESTs
2079	15701	NM_080581	j,m,y,z	binding cassette, sub-family C (CFTR/MRP), member 3 (Abcc3), mRNA. Length = 5174	Rattus norvegicus mRNA for multidrug resistance-associated protein (MRP)- like protein-2 (MLP-2), complete cds
2079	20105	NM_080581	aa	Rattus norvegicus ATP- binding cassette, sub-family C (CFTR/MRP), member 3 (Abcc3), mRNA. Length = 5174	ESTs
					Rattus norvegicus mRNA for alpha-
2080	1757	NM_080585	d	Length = 1505 Rattus norvegicus Neuroblastoma RAS viral (v- ras) oncogene homolog (Nras), mRNA. Length = 3350	R.norvegicus N-ras gene for p21
2082	7108	NM_080778	у	Rattus norvegicus nuclear receptor subfamily 2, group F, member 2 (Nr2f2), mRNA. Length = 1572	Rattus norvegicus ovalbumin upstream promoter beta nuclear receptor rCOUPb mRNA, complete cds
2083	132	NM_080782	k	Rattus norvegicus cyclin- dependent kinase inhibitor 1A (P21) (Cdkn1a), mRNA. Length = 495	Rattus norvegicus p21 (WAF1) mRNA, complete cds
2083	133	NM_080782	ı	Rattus norvegicus cyclin- dependent kinase inhibitor 1A (P21) (Cdkn1a), mRNA. Length = 495	Rattus norvegicus p21 (WAF1) mRNA, complete cds
2084	20122	រា NM_080887	General	Rattus norvegicus thioredoxin-like (32kD) (Txnl), mRNA. Length = 1061	ESTs, Highly similar to thioredoxin- related protein [M.musculus]
2085	6143	NM_080892	е	mRNA. Length = 1685	ESTs, Moderately similar to selenium- binding protein [H.sapiens]
2086	9952	NM_080902	h	Rattus norvegicus hypoxia induced gene 1 (Hig1), mRNA. Length = 355 Rattus norvegicus	ESTs, Moderately similar to AF077034 1 HSPC010 [H.sapiens]
2087	17546	NM_130401	b	membrane-associated protein 17 (Map17), mRNA. Length = 816	ESTs, Moderately similar to DD96 homolog [R.norvegicus]

TABLE 1: 8	XAIXIXXA <u>*</u>				Atty, Docket No. 44921-5059W0 Doc. No. 1793397.
Sequence ID No.	relitine o	Genbank Axel Ref. Seg ID	Model.	Cone Name	Unigene Cluster Title
				Rattus norvegicus coronin, actin binding protein 1A (Coro1a), mRNA. Length =	ESTs, Weakly similar to coronin-like
2088	21695	NM_130411	c,x	1386 Rattus norvegicus annexin	protein [R.norvegicus]
2089	21391	NM_130416	x,General	, , , , ,	ESTs, Weakly similar to ANX4 RAT ANNEXIN IV [R.norvegicus]
2090	20694	NM 130430	General	proteasome (prosome, macropain) 26S subunit, non- ATPase, 9 (Psmd9), mRNA. Length = 1448	EST
2090	20094	130430	Concra	Rattus norvegicus proteasome (prosome, macropain) 26S subunit, non- ATPase, 9 (Psmd9), mRNA.	
2090	19818	NM_130430	СС	Length = 1448	EST
2000	40040	NN 120420		Rattus norvegicus proteasome (prosome, macropain) 26S subunit, non- ATPase, 9 (Psmd9), mRNA.	mitochondrial H+-ATP synthase alpha
2090	18810	NM_130430	e,s	Length = 1448	subunit
				Rattus norvegicus acetyl- Coenzyme A acyltransferase 2 (mitochondrial 3-oxoacyl- Coenzyme A thiolase) (Acaa2), mRNA. Length =	
2091	18293	NM_130433	g	1619	Rat mRNA for 3-oxoacyl-CoA thiolase
2092	25064 3244	S45392 S63519	a,n u		ESTs
2094	25501	S63521	q		
2095	16248	S68135	h		Rat brain glucose-transporter protein mRNA, complete cds
					ESTs, Weakly similar to HS9B RAT HEAT SHOCK PROTEIN HSP 90-
2096	18647	S69316	q		BETA [R.norvegicus] ESTs, Weakly similar to ABD4 MOUSE ATP-BINDING CASSETTE,
0007	04054	074057			SUB-FAMILY D, MEMBER 4 [M.musculus]
2097 2098	24351 25066	S74257 S75280	d		[M.Musculus]
		\$76054	j,l,m,x,y,G		ESTs, Highly similar to K2C8 RAT KERATIN, TYPE II CYTOSKELETAL 8 [R.norvegicus]
2100	1460 25539	S76742	eneral v		o [ix.norvegicus]
2101	16400	\$76779	С		Rat apolipoprotein e mrna ESTs. Highly similar to MLES RAT
					MYOSIN LIGHT CHAIN ALKALI, SMOOTH-MUSCLE ISOFORM
2102 2103	24469 25545	S77858 S77900	n k,s		[R.norvegicus]
2103	21583	S77900 S77900	k		ESTs
2104	10260	S81497	s		ESTs
2105	3609	S82579	k	histamine N- methyltransferase	histamine N-methyltransferase Rattus norvegicus clone 15 polymeric
2106	111	U02506	u		immunoglobulin receptor mRNA, 3'UTR microsatellite repeats Rattus norvegicus Sprague Dawley
2107	14959	U03390	a,q, General		protein kinase C receptor mRNA, complete cds
	2010	U05675	b,x,bb		Rattus norvegicus Sprague-Dawley fibrinogen B beta chain mRNA, complete cds
2109	170110	11105675	IN Y PP	1	Icomplete cas

TABLE 1:	SUMMARY:				Any, Doeket No. 44921-5039WC Doe, No. 1793897.1
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Sequence ID No.	ldeniiiler	GenBank Acc/ Ref. Seq ID	Model Code	Gene Namo	Unigeno Civeter Tilo
2112	1583	U07201	s.General	Asparagine synthetase	Asparagine synthetase
					Rattus norvegicus New England Deaconess transcription factor mRNA,
2113	627	U09229	h		Partial cds Rattus norvegicus interferon inducible protein 10 (IP-10) mRNA, complete
2114	809	U17035	General		cds
				mini chromosome maintenance deficient 6 (S.	mini chromosome maintenance
2115	16675	U17565	k,x,bb	cerevisiae)	deficient 6 (S. cerevisiae)
2116	25587	U20110	r		Rattus norvegicus nuclear receptor
2117	90	U20796	r		Rev-ErbA-beta mRNA, partial cds
2118	25589	U21718	h,aa		
2119	22196	U21719	h		ESTs
	15110				Rattus norvegicus RNA helicase with arginine-serine-rich domain mRNA,
2120	17118	U25746	s		complete cds Rattus norvegicus UDP-
2121	1537	U27518	g,h,n		glucuronosyltransferase mRNA, complete cds
2121	1007	02.010	9,,,,,	· · · · · · · · · · · · · · · · · · ·	Rattus norvegicus Na+/Pi
2122	1558	U28504	bb		cotransporter-1 mRNA, complete cds Rattus norvegicus B/K protein mRNA,
2123	16193	U30831	n		complete cds
2124	17480	U31598	z		R.norvegicus mRNA for RT1.Ma
0405	40000	1100500	C		Rattus norvegicus retinol dehydrogenase type II mRNA, complete cds
2125 2126	18302 25599	U33500 U34897	General v		complete cas
2120	20000	304001	/		Rattus norvegicus GTP-binding
2127	1394	U37099	h		protein (rab 3C) mRNA, complete cds EST, Weakly similar to actin-filament
2128	244	U38376	n		binding protein Frabin [R.norvegicus],Rattus norvegicus cytosolic phospholipase A2 mRNA, complete cds
2120	244	038376	11		Rattus norvegicus Cys2/His2 zinc
2129	1623	U41164	ħ		finger protein (rKr1) mRNA, complete cds
2130	15851	U42719	f,t,x, General	Complement component 4	Complement component 4
2130	13031	042718	General	Complement component 4	Rattus norvegicus apoptosis-
2131	17886	U47315	s,z		regulating basic protein mRNA, complete cds
			i,t,	estrogen-responsive uterine	
2132	21654	U53184	General	transcript	Rattus norvegicus FceRl gamma-
2133	1439	U57391	l _w		chain interacting protein SH2-B (SH2-B) mRNA, complete cds
				solute carrier family 16	solute carrier family 16
2134	725	U62316	bb	(monocarboxylic acid transporters), mem	(monocarboxylic acid transporters), member 7
2137	2153	U75404	b,cc, General		ESTs
					Rattus norvegicus cell adhesion regulator (CAR1) mRNA, complete cds
2139	4956 4477	U76714 U77829	j,y I,m		ESTs
2141	21703	U82591	z		Rattus norvegicus RCL (Rcl) mRNA, complete cds
					Rattus norvegicus putative cell
2142 2143	977 23282	U89744 U90725	s h	lipoprotein-binding protein	surface antigen mRNA, complete cds lipoprotein-binding protein
2144	22005	U96490	m		Rattus norvegicus liver mRNA, complete cds

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TABLE 1: 8					AMy. Docker No. 44921-5039W0 Doc. No. 1793397.1
Sequence ID No.	ldenMier	GenBank Acel Ref. Seque	Model Model	Gene Name	Unicene Circler Mile
The state of the s		Sourcement of Sections - Address (All Conference)	en manudament ext. 1 Ag	Aldolase B, fructose-	
2146	819	X02284	j,z	biphosphate	Aldolase B, fructose-biphosphate
2147	818	X02291	e,j,z	Aldolase B, fructose- biphosphate	Aldolase B, fructose-biphosphate
				glutathione S-transferase, pi	
2148	20818	X02904	n,q	2	glutathione S-transferase, pi 2
2149	16401	X04979	С	Pyruvate kinase, liver and	Rat apolipoprotein e mrna
2150	20513	X05684	o,r	RBC	Pyruvate kinase, liver and RBC
2151	25084 672	X06769 X13722	cc h		Rat mRNA for LDL-receptor
2152 2153	25675	X13722 X14181	n		Rat IIIRNA for EDE-receptor
2153	20810	X14181	n,q,w		ESTs, Highly similar to 60S RIBOSOMAL PROTEIN L18A [R.norvegicus]
			11,q,w		ESTs, Highly similar to RL26 RAT 60S RIBOSOMAL PROTEIN L26
2154	18541	X14671	<u>y</u>		[R.norvegicus]
2155	25679	X15013	q		ESTs, Highly similar to RL7A_HUMAN 60S RIBOSOMAL PROTEIN L7A
2155	19244	X15013	c,q,w		[R.norvegicus] ESTs, Highly similar to RS16_HUMAN 40S RIBOSOMAL PROTEIN S1
2156	15626	X17665	а	phospholipase A2, group IIA	[R.norvegicus] phospholipase A2, group IIA
2157	1893	X51529	t	(platelets, synovial fluid)	(platelets, synovial fluid)
2158	25686	X51536	bb		
0.450	40040	V54500			ESTs, Highly similar to RS3 MOUSE 40S RIBOSOMAL PROTEIN S3
2158	10819	X51536	aa,bb		[R.norvegicus] ESTs, Highly similar to RL9 RAT 60S RIBOSOMAL PROTEIN L9
2159	18250	X51706	a,q,w	ribosomal protein L9	[R.norvegicus] ESTs, Highly similar to RS19 RAT
2160	20872	X51707	а	ribosomal protein S19	40S RIBOSOMAL PROTEIN S19 [R.norvegicus]
2161	516	X52711	c	,	Rat mRNA for Mx1 protein
2162	25689	X52815	g		•
2163	20427	X53378	w		Rattus norvegicus ribosomal protein S13 (RPS13) mRNA, 3' end
2464	40000	V52504	Canaral		ESTs, Highly similar to 60S RIBOSOMAL PROTEIN L12
2164	18606	X53504	General d,u,		[R.norvegicus] Rat mRNA for preprocathepsin D (EC
2165	1463	X54467	General		3.4.23.5) ESTs, Highly similar to 60S ACIDIC
2166	24577	X55153	a,v		RIBOSOMAL PROTEIN P2 [R.norvegicus]
2167	10344	X57405	j,m	Drosophila Notch homolog 1	LOCUS NOTCH HOMOLOG PROTEIN 1 PRECURSOR [R.norvegicus]
2168	15106	X57529	g,n,q		ESTs, Highly similar to RS18_HUMAN 40S RIBOSOMAL PROTEIN S18 [R.norvegicus]
2169	5667	X58200	q,bb	ribosomal protein L23	
					ESTs, Highly similar to RL23_HUMAN 60S RIBOSOMAL PROTEIN L23
2169	18611	X58200	a,v	ribosomal protein L23	[R.norvegicus] R.norvegicus ASI mRNA for mammalian equivalent of bacterial
2170	17175	X58389	w		large ribosomal subunit protein L22
2171	25702	X58465	w		
2171	10109	X58465	c,q	Ribosomal protein S5	Ribosomal protein S5
2172	25705	X59375	c,i,aa, General		
				L	

TABLE 1: 8					Atty. Docket No. 44921-5089W0
	No.				Doc. No. 1793697.1
Sequence ID No.	I CANADA	Conbank Acc Roi. Sog ID	Model Gode	Gene Name	Unigene Gluster Tille
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2173 2174	25709 18354	X59737 X59859	u General	decorin	decorin
2174	18355	X59859	t	decorin	decorin
	10000	7.03003	<u> </u>	Tubbonii Tub	Rattus norvegicus interferon-inducible
2175	21657	X61381	General		protein variant 10 mRNA, complete cds
			bb,		
2176	25718	X62145	General	ribosomal protein L8	ESTs, Highly similar to RL8_HUMAN
2176	15875	X62145	a,q,v	ribosomal protein L8	60S RIBOSOMAL PROTEIN L [R.norvegicus]
	10010				ESTs, Highly similar to RL3 RAT 60S RIBOSOMAL PROTEIN L3
2177	13646	X62166	bb		[R.norvegicus]
2178	25721	X62325	P		ESTs, Highly similar to HIGH
2179	16012	X62875	m,s,z		MOBILITY GROUP PROTEIN HMG-Y [M.musculus]
2180	25730	X63369	CC		
2181	25089	X63594	General		
			cc,	1	
2181	25090	X63594	General		ESTs, Highly similar to RL2B_HUMAN
0400	00044	V05000	_		60S RIBOSOMAL PROTEIN L23A
2182 2183	20844	X65228 X65296	n,w	carboxylesterase 1	[R.norvegicus] carboxylesterase 1
2184	25736	X68782	j.y c	Carboxylesterase i	Carboxylesterase i
2185	16426	X70369	c	procollagen, type III, alpha 1	procollagen, type III, alpha 1
2186	16300	X70706	u	plastin 3 (T-isoform)	plastin 3 (T-isoform)
2187	24232	X75207	c	cyclin D1	cyclin D1
2188	16272	X76456	n,p		R.norvegicus (Sprague Dawley) alpha albumin gene
2189	25741	X76489	u		
2190	23302	X78949	h	prolyl 4-hydroxylase alpha subunit	prolyl 4-hydroxylase alpha subunit
2191	25747	X81448	General		
0400	04445	V04440			ESTs, Highly similar to K1CS RAT KERATIN, TYPE I CYTOSKELETAL 19 [R.norvegicus]
2192 2193	24115 25754	X81449 X89696	g g		19 [K.Horvegicus]
2194	25097	X90642	y,z		
			cc, General		ESTS, Highly similar to RADIATION- INDUCIBLE IMMEDIATE-EARLY
2195	12978	X96437			GENE IEX-1 [M.musculus] Rattus norvegicus mRNA Best5
2197	4594	Y07704	C C		protein
2198	25777	Y08355	g.p, General	oxidative stress induced	oxidative stress induced Rattus norvegicus mRNA for putative
2199	15986	Y09945	bb, General		integral membrane transport protein (UST1r)
2200	20890	Y13275	k		Rattus norvegicus mRNA for D6.1A protein
2201	21914	Y13336	d		Rattus norvegicus DAD-1 gene
					R.norvegicus mRNA encoding 45kDa protein which binds to heymann
2202	406	Z11995	o,General		nephritis antigen gp330
2203	18352	Z12298	t	decorin	decorin
2204	17481	Z49761	k		R.norvegicus mRNA for RT1.Ma ESTs, Moderately similar to T17342 hypothetical protein DKFZp586K1924.1 [H.sapiens],Heat
2205	8664	Z75029	r,v	Heat shock protein 70-1	shock protein 70-1
2206	2459	AA964755	СС		ESTs
2207	23830	AA956638	aa		ESTs
2208	6100	X73524	х	desmin	desmin

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TABLE 1: 8				11/4/2	Ally, Docket No. 44921-5039\/
		e de la composição de l	1000 b. 1425		Doc. No. 1793397.
Sequence ID No.	ldeniller	Consilk Acc		Cono Namo	Unigane Cluster Title
2209	439	Z22607	w	Bone morphogenetic protein 4	Bone morphogenetic protein 4
2210	8665	AI071965	v	Heat shock protein 70-1	ESTs, Moderately similar to T17342 hypothetical protein DKFZp586K1924.1 [H.sapiens],Heat shock protein 70-1
2211	155	U32681	t	crp-ductin	crp-ductin
2212	19252	AA892041	s	HMm:peroxiredoxin 5	Rattus norvegicus mRNA for thiol- specific antioxidant protein (1-Cys peroxiredoxin)
2213	15582	AI232320	q		Rat mitochondrial 3-hydroxy-3- methylglutaryl-CoA synthase mRNA, complete cds
2214	17541	M26125	n	Epoxide hydrolase 1 (microsomal xenobiotic hydrolase)	Epoxide hydrolase 1 (microsomal xenobiotic hydrolase)
2215	18609	M30689	i		Rat Ly6-B antigen mRNA, complete cds
2216	6262	AI177125	g		ESTs
2217	23859	AI072161	f		ESTs
2218	21011	H32189	е	Glutathione-S-transferase, mu type 2 (Yb2)	Glutathione-S-transferase, mu type 2 (Yb2)
2220	2572	AI177143	b		ESTs
2221	25419	M22922	а		

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College to the secretary of the second secon	ZYZWWAYYS			Atty, Docks No. 44921-5089Wo Doc, No. 1798897.1
Sequence ID No.	lden ii ier	Genfenk Acel Ref. Seq. ID No.	Modal Goda	Pathways 442
1	6949	AA012785	q	
2	25098	AA108277	h,v	
3	17312	AA108308	r	
4	16882	AA684537	0	
5	6049	AA685178	у	
6	4426	AA685974	I,m	
7	21815	AA686423	g :	
8	1600 1599	AA686470 AA686470		
9	21997	AA799325	u	
10	18396	AA799330	v	
11	6581	AA799412	f,l	
12	16538	AA799449	k	
13	23294	AA799472	u	
14	18290	AA799497	r	
15	18981	AA799523	е	
16	20843	AA799545	h	
17	16993	AA799560	b	
18	16576	AA799570	d	
19	18361	AA799591	<u> -</u>	
20	17712	AA799598	Z	
22	18346	AA799718 AA799726	f	
23 24	8768 11687	AA799732	w	
25	18349	AA799744	u	
26	17494	AA799751	n	
27	18360	AA799771	General	
28	18880	AA799801	w	
29	20998	AA799803	z	
30	21006	AA799861	С	
31	15011	AA799893	General	
32	20811	AA799899	а	
33	23202	AA799971	General	
34	4832	AA800190	b	
35	21656	AA800202	d	
36 37	18433 6386	AA800218 AA800235	j,y,z u	
38	18442	AA800258	h,k	
39	21092	AA800380	y	
40	17325	AA800587	General	<u> </u>
41	13930	AA800613	cc,General	
42	21372	AA800693	V	
42	21373	AA800693	s	
43	18161	AA800701	k	
44	6595	AA800753	w	
45	13348	AA800928	General	
46	23115	AA801165	O,y	
47	12399	AA801307	General	
48 49	7543 24237	AA801395 AA817726	General t,General	
50	11215	AA817726 AA817921	t,General o	
51	5985	AA818005	g	
52	11338	AA818016	X	
53	2845	AA818026	k,General	
54	16756	AA818089	i,k,General	
55	17771	AA818224	e,g,p,General	
56	6522	AA818261	g,m	
57	5924	AA818359	У	
58	7806	AA818421	b,aa	
59	8237	AA818512	V	
60	17434	AA818574	h Canaral	
61 62	8728	AA818615	General	
02	6054	AA818658	b,v,cc,General	

vable2. F				(Atiy: Podkatiko. 44921-5089) Doc. Ko. 179899
Sequence D No. ''	(dentifier	Canbank Acel Ref. Seq. ID No. 1	Model Gode	(Pathways)
53	11590	AA818721]d	
64	4291	AA818741	q,General	
65	4330	AA818747	o,General	
36	19723	AA818761	v,General_	
57	13684	AA818770	h,j,l,m	
68	6322	AA818801	Conoral	
69 70	7690 4952	AA818875 AA818907	General q,General	
70 71	6094	AA818911	t	
72	10985	AA818998	o,General	
73	6120	AA819008	t	
74	2586	AA819081	С	
76	6438	AA819269	0	
77	24721	AA819306	d,w	
78	6250	AA819376	o,y	
30	6281	AA819517	j	
31	10141	AA819526	j	
82	6551	AA819558	t	
83	6723	AA819653	r	
84	14958	AA819744	aa 	
35	19433	AA819776	V	
86 87	6204 22820	AA819889 AA848315	aa General	Purine metabolism
88	6614	AA848389	bb	Purine metabolism
39	21125	AA848437	General	
90	23504	AA848496	q	
91	18532	AA848675	9	
92	21140	AA848738	C	
93	16128	AA848807	0	
94	22923	AA848929	g	
95	17339	AA849497	General	
96	11727	AA849518	1	
97	21275	AA849796	i,I,m,General	
98	16678	AA849827	aa	
99	8515	AA849917	е	
100	18447	AA849939	General	
101	12130	AA850037	p	
102	23981	AA850040	x,aa	
103	13615 2637	AA850364 AA850893	t -	AP-22
105 106	22093	AA850893 AA850909	d d	
107	21766	AA850916	c	
108	2847	AA850919	w	
109	12162	AA850975	h	
110	9514	AA850978	General	
111	3924	AA851017	e,q	
111	3925	AA851017	o,General	
112	4490	AA851184	a,k	
113	19187	AA851230	General	
114	19189	AA851237	С	
115	15386	AA851241	m	
116	21462	AA851261	g,I,General	
117	21471	AA851343	General	
40	1,,,,,,			Oxidative phosphorylation, Ubiquinone
118	16902	AA851379	p	biosynthesis
119	23376	AA851392	i,x	
119	23377	AA851392	Conoral	
120	13349	AA851417	General	
21 22	21527 4048	AA851733 AA851814	r,u i,o,u,General	
123	10561	AA851871	bb	· · · · · · · · · · · · · · · · · · ·
124	17411	AA858621	j,y	
125	1801	AA858636	k,s,x,bb	

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TABUE 2: P	ATHWAYS			ATTY, Docket No. 44921-5039W0 Doc. No. 1793397.1
Ecquence	ldendder	Cenbenk Acel Ref. Seq. 10 No.	Model Gode	
126	18350	AA858674	р	
127	19484	AA858693	е	
128	6360	AA858696	d	
129 130	17334 6380	AA858704 AA858758	p q	
131	13219	AA858759	a	
132	6384	AA858788	I,m,General	
134	13412	AA858830	р	
135	7279	AA858892	f	
136	18217	AA858930	<u>t</u>	Al
407	5007	A A 0 E 0 0 E 2	v.General	Alanine and aspartate metabolism, Aminoacyl-tRNA biosynthesis
137 138	5867 14479	AA858953 AA858969	r,General_	metabolism, Aminoacyi-trina biosynthesis
139	6431	AA859085	t	
140	17361	AA859114	o,General	
141	21025	AA859241	General	
142	10076	AA859271	С	
143	21791	AA859333	k	
144	16314	AA859348	cc,General	
145 146	18862 15059	AA859520 AA859545	r	
147	19894	AA859581	s	
148	14353	AA859585	h	
149	16318	AA859648	h	
150	17316	AA859652	General	
151	19067	AA859663	n,q	
152	22406	AA859680	n	
153	20599	AA859690	x	
154 155	14261 14138	AA859693 AA859700	u v	Porphyrin and chlorophyll metabolism
155	14139	AA859700	v	Porphyrin and chlorophyll metabolism
157	22374	AA859804	i	· · · · · · · · · · · · · · · · · · ·
158	22385	AA859805	b,k	
159	22773	AA859885	n	
160	22816	AA859898	k,x,z	
161	11891	AA859926	X	
162	23070	AA859942 AA859948	k	
163 164	23121	AA859946 AA859954	cc.General	****
165	18468	AA859966	aa	
166	23336	AA859981	q	Inositol phosphate metabolism
167	4222	AA860024	a,bb	
168	13974	AA860030	u,x,General	
169	7090	AA860039	x	
170	23769	AA860055	k,x	
171 172	16323 4462	AA866240 AA866264	w General	
173	15884	AA866276	k	
	1,000+	7 0 10002 7 0	<u> </u>	Phenylalanine metabolism, Tyrosine
174	17742	AA866302	с,у	metabolism
175	16333	AA866414	a,h	
176	18918	AA866444	p,q	
177	16853	AA866454	j,l,m,y,z	
178	18995	AA866459 AA866482	h,m	
179 180	16013 26036	AA874849	s r	
181	16059	AA874857	h	<u> </u>
182	16069	AA874873	r	· · · · · · · · · · · · · · · · · · ·
183	21633	AA874951	f	
184	16192	AA874995	w	
185	16254	AA875025	ji	
186	16312	AA875032	cc,General	
187	20701	AA875097	b	I

TABLE 2: P	ATHWAYS			449241-5034 Doc. No. 17933
Sequence D No.	ldentifier	ConBank Ace/ Ref. Seq. ID No.	Model Code	Pathways.
188	16416	AA875098	bb	
89	16419	AA875102	bb	<u> </u>
190	15313	AA875126	I,m,General	
191	10936	AA875146	w	
192	18084	AA875186	h	
193	15371	AA875205	u	
194	15401	AA875257	x,z	
195	15410	AA875268	p,s	Oxidative phosphorylation, Ubiquinone biosynthesis
96	15420	AA875286	f	
197	15446	AA875327	s,w	
198	7936	AA875495	b,General	
199	17314	AA875509	i,l,m	
200	24472	AA875523	k	
201	15587	AA875577	ji	
202	15617	AA875620	General	
202	15618	AA875620	General	
203	5384	AA891041	f,cc,General	
204	24814	AA891209	f,p	
205	21930	AA891322	d	
206	17225	AA891553	h	
207	7522	AA891571	j,m	
208	9071	AA891578	b	
209	19321	AA891666	u	
210	17693	AA891737	j,l,m,n,y,z	
211	17256	AA891739	General	
213	18269	AA891769	General	
214	9905	AA891774	s,bb,General	
215	17061	AA891812	d	
216	7050	AA891824	h	
217	4463	AA891831	General	
218	14289	AA891838	i	
219	20523	AA891842	r,cc	
220	17779	AA891914	g,s,z	
221	17438	AA891943	General	
222	22862	AA891944	р	
223	1159	AA891949	e,z	
224	4473	AA891965	General	
225	6362	AA892053	f,j,l,m	
226	9037	AA892066	У	
227	19469	AA892112	General	
228	14595	AA892128	o,t,v	
229	16527	AA892154	cc	
230	4482	AA892173	bb	
231	20917	AA892238	h	
232	2357	AA892268	d	
233	18183	AA892271	h	
234	6523	AA892299	d	
236	13647	AA892367	a	
237	3473	AA892378	v	
238	17682	AA892382	j,p,s,x, General	
				Carbon fixation, Fructose and mannose metabolism, Glycolysis / Gluconeogenesis, Inositol
239	820	AA892395	g,s	metabolism,Pentose phosphate cycle
240	14754	AA892414	u	
241	17439	AA892446	f	
242	16469	AA892462	р	
243	13609	AA892468	i,General	
243	13610	AA892468	n,v,General	
244	9254	AA892470	n,u	
245	11991	AA892483	s	
246	1522	AA892486	f	

MARUE 28 F				<u>Doc. No. 1793397</u>
Boguence D No.	ldeniiier	ConBank Acel Ref. Seq. ID No.	Model Gode	Pathways ************************************
247	11994	AA892507	aa	
248	23888	AA892520	w	
248	23889	AA892520	h	
249	8599	AA892522	p	
250	15154	AA892532	p r	
251 252	17468 11203	AA892545 AA892554	f,h	
253	18906	AA892561	a.bb.General	
254	19327	AA892562	f,j,y,z	
255	18274	AA892572	p	
256	4512	AA892578	СС	
257	15876	AA892582	w	
258	19085	AA892598	General	
258	19086	AA892598	General	
259	20065	AA892647	-	<u> </u>
260	20088	AA892666	a,n	
261 262	23783 17549	AA892773 AA892776	n f,z	
263	13542	AA892776 AA892798	b	
203	10042	AA032730	<u> </u>	Glyoxylate and dicarboxylate
264	22537	AA892799	General	metabolism,Pyruvate metabolism
		7 (302/30		Glyoxylate and dicarboxylate
264	22539	AA892799	v	metabolism,Pyruvate metabolism
-				Glyoxylate and dicarboxylate
264	22538	AA892799	General	metabolism,Pyruvate metabolism
265	6951	AA892820	h	
266	23322	AA892821	j,z	
267	17923	AA892843	f	
268	22871	AA892859	m 	
269 270	9053 16482	AA892861 AA892940	p,v,General w	
271	12020	AA893035	i,y	
272	3863	AA893060	General	
273	13332	AA893080	i,General	
274	21305	AA893082	General	
275	16591	AA893191	j,z	
276	17447	AA893192	General	
277	3876	AA893205	n	
278	3878	AA893230	General	
279	20986	AA893242	9	Fatty acid metabolism
280	16168	AA893280	i,z,General	
281	3886	AA893289	j,m,y	-
282 283	15209 17800	AA893327 AA893436	cc	
284	17836	AA893626	h	
285	9084	AA893717	x	
286	22731	AA893743	d	
	1	1.2.2.2		Aminoacyl-tRNA biosynthesis, Glycine,
287	12031	AA893860	v	serine and threonine metabolism
288	17897	AA893905	k	
289	3447	AA893982	d	
90	22583	AA894009	n	
91	10540	AA894027]	
292	4569	AA894059	X	
293 294	18419	AA894130	d	
294 295	17336 19120	AA894297 AA894318	f,j	· · · · · · · · · · · · · · · · · · ·
.95 .96	19762	AA899113	'.J	
297	18286	AA899219	u	
298	22051	AA899498	w	
298	22052	AA899498	q	
299	21628	AA899563	aa	
300	4262	AA899590	i	<u> </u>

TABUE 2: F	ALLINA ALLINA		<u> </u>	Aliy: Docto: No. 44921-5039W0 Doc. No. 1793397.1
Sequence		Cenfenk Acel Ref. Seg. ID No.	Model Gode	Pathways 1
301	4661	AA899709	t,General	
302	21354	AA899721	q .	
303	17905	AA899762	General	
304	15231	AA899840	r	
305	23778	AA899854	c,k,x	
306	22060	AA899898	b v.General	
307 308	9114 8988	AA899951	v,General	
309	11841	AA900148 AA900247	ľ	
310	4725	AA900290	cc	
311	4747	AA900465	General	
312	20988	AA900562	0	
313	3822	AA900863	b,g,General	
315	12420	AA901017	b	
316	4849	AA901155	s	
317	3959	AA901338	General	
318	22846	AA923982	a,d	
319	4895	AA923999	k Constal	
320	21546	AA924188	cc,General	
321	24192	AA924210	n,General g,I,General	
322	4933	AA924301 AA924405	I,General	
323 324	4944 4948	AA924405 AA924428	r,General	
325	4949	AA924426 AA924432	General	
326	18891	AA924598	e	
327	22540	AA924630	v,General	Glyoxylate and dicarboxylate metabolism,Pyruvate metabolism
327	22541	AA924630	General	Glyoxylate and dicarboxylate metabolism, Pyruvate metabolism
328	14759	AA924766	k	metabolism, yruvate metabolism
329	23123	AA924794	x	
330	4067	AA924813	g,p	
331	2888	AA924902	r,General	
332	18130	AA924964	d	
333	23141	AA925019	r	
334	23195	AA925026	General	
335	21458	AA925049	f,aa,General	
336	5073	AA925061	m	****
337	14790	AA925087	o,General	
338	5089	AA925126	g la Comerci	
339	23261 17363	AA925145 AA925150	k,General a	
340 341	23448	AA925167	ı	
342	23159	AA925318	e	
343	21500	AA925353	k	
344	22479	AA925418	t	
345	21151	AA925539	b	
346	16944	AA925541	f	
346	16945	AA925541	t	
347	17514	AA925554	bb	Oxidative phosphorylation
348	5183	AA925662	i,General	
349	23189	AA925844	r	
350	23190	AA925863	aa	
351	5252	AA926051	General	
352	22967 17157	AA926080 AA926129	h,cc b	
353 354	13411	AA926196	u,General	
355	5295	AA926196 AA926247	General	
356	22928	AA926262	General	
357	8948	AA926316	r	
358	21798	AA926365	aa	
359	9942	AA942697	s	
360	6039	AA942716	x,General	

vabue 2: F	ATHWAYS			Aviy, Docket No. 44921-5039W Doc. No. 1798397
ID No. : : : : Sequence :	lieniliter	Cenbenk Acel Ref. Seq. ID No.		Pathways : ##
361	11174	AA942745	g,o,w	
362	23005	AA942770	g	
363	21318	AA942774	General	
364	6615	AA942889	V	
365	6691	AA943028	С	
366	22142	AA943066	p v,General	<u> </u>
367 368	21993 9061	AA943149 AA943508	General	
369	24390	AA943531	b,j,n,y	
370	13976	AA943532	f,s,x	
371	22248	AA943537	cc,General	
372	22257	AA943558	m	
373	12673	AA943773	u,cc,General	
374	13641	AA944154	u	
375	2658	AA944155	f	
376	12770	AA944161	d	
377	20903	AA944180	i,x	
378	13507	AA944244	V Canada	
379	15596	AA944353	General	
380	22681	AA944413	i,v,cc,General General	
381 382	6711 14763	AA944439 AA944481	i,q,General	
383	22466	AA944605	h	
384	12301	AA944727	b	
304	12301	77344121		Purine metabolism,Pyrimidine
385	7023	AA944792	d,m,aa	metabolism,RNA polymerase
386	22536	AA944803	bb	
387	22501	AA944811	g,l	
388	23967	AA944831	S	
389	26084	AA944922	i	
390	11974	AA944958	General	
391	22547	AA944970	aa	
392	22554	AA945076	z,General	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
393	14352	AA945181	General	
395	1798	AA945569	General	
396 397	22050 19731	AA945604 AA945615	i,aa d,o	
398	22612	AA945624	a,General	
399	22618	AA945656	aa	
400	11871	AA945679	v	·
401	22656	AA945818	General	
402	6720	AA945828	р	
403	22351	AA945867	m	
404	22665	AA945877	f	
405	24243	AA945950	b	
406	22689	AA945962	General	
107	22692	AA945986	d	
108	22696	AA945996	c,General	
108	22697	AA945996	c,o	
409	22658	AA945998	w	Ovidative phosphanilation
410	20832	AA946040 AA946046	S	Oxidative phosphorylation
411 412	18337 825	AA946108	General General	
113	8639	AA946221	e,cc,General	
¥14	23237	AA946224	f	
415	15600	AA946250	o,aa	
416	19387	AA946275	t	
417	6351	AA946344	d	
418	22057	AA946348	е	
419	22069	AA946349	aa	
420	13962	AA946351	General	
421	18280	AA946361	g	
422	18944	AA946391	V	

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TABLE 28 P			Maria di	Affiy, Docket No. 44921-503900 Doc. No. 1793397
Sequence ID No.	ldentifer	ConBank Acel Ref. Soq. ID'No.	Model Gode	Pathways ***
424	21410	AA946408	t	
425	643	AA946439	o,y	
426	20736	AA946443	x	
427 428	21878 21947	AA946448 AA946451	r bb	
420 429	17499	AA946467	General	
430	1809	AA946503	x.General	
431	23360	AA955104	f	
432	23471	AA955162	General	
433	9452	AA955206	b,General	
434	23512	AA955282	General	
435	22596	AA955298	General	
436	23283	AA955391	h Canasal	
437	23546 12404	AA955393 AA955408	General b	
438 439	23626	AA955408	aa	
439 441	17540	AA955914	bb	
442	24277	AA955962	General	
443	19939	AA955980	General	
444	24000	AA956005	i	
445	11050	AA956164	s,v	
446	498	AA956278	a,General	
447	23409	AA956294	q	
449	23773	AA956476	f,x	
450	23799	AA956530	d	
451	23800	AA956534	aa	
452 453	23834 16425	AA956659 AA956688	cc,General f,x	
453 454	23847	AA956723	s	
455	23852	AA956746	j,l,m,z	
456	5989	AA956907	g,s	
456	5990	AA956907	General	
457	23957	AA957123	u,General	
458	22357	AA957264	General	
			g,l,m,p,v,cc,	
459	23314	AA957270	General	
460	23995	AA957292	a,b	
				Aminoacyl-tRNA biosynthesis, Glycine,
461	2702	AA957307	General	serine and threonine metabolism
462 463	24040	AA957422	m	
464	12478 21306	AA957554 AA957811	v	
465	24183	AA957889	t	
466	24178	AA957905	d	10
467	17034	AA963071	е	
468	24053	AA963092	General	
469	2767	AA963201	0	
470	2022	AA963259	g	
471	2126	AA963488	d	
472	24246	AA963703	General	
473 474	2195 19370	AA963746 AA963797	General	
474 475	2282	AA964147	e	
476	2284	AA964152	x	
478	2350	AA964368	g,General	
479	18830	AA964496	aa	
480	2392	AA964541	b	
481	2395	AA964554	General	
482	2410	AA964589	i,aa	
483	19145	AA964613	t	
484	2424	AA964617	g Consent	
485	3107 2457	AA964687 AA964752	General q,t	

MABLE 28 P	ATHWAYS	were provided a section		(X i ly), Dogital No. 449911-5039VV
				Dog. No. 17/98397
Sequence ID No. :	ldeniliter	CenBank Acci Ref. Seq. ID No.	Model Code	Pathways
487	6778	AA964763	b	
489	2468	AA964807		
490	2469	AA964814	w	Glutamate metabolism, Glutathione metabolism
491	12561	AA964815	General	
492	2326	AA964892	aa	
493	21339	AA964962	General	
494	21390	AA964988	General	
495	12569	AA965023	g	
496	2583	AA965166	bb	
497	15885	AA965207	r	
499	2905	AA996727	b,l,m,u,General	
500	2915	AA996782	u,bb	
501 502	2920 19525	AA996813 AA996856	d aa,General	
502 503	2984	AA990000 AA997015	c c	
504	2986	AA997015 AA997028	General	
505	3145	AA997020 AA997237	General	
506	19249	AA997342	m	
507	16883	AA997345	General	
508	12598	AA997362	s	
509	3470	AA997374	р	
510	3180	AA997425	t	
511	3245	AA997608	General	
512	3020	AA997656	t	
513	3269	AA997800	x,aa	
514	3288	AA997877	<u> </u> †	
515	23992	AA998164	k,x	
516	17470	AA998264	b General	
517 518	3773 19623	AA998356 AA998422	General	-
519	3572	AA998516	x	
520	2782	AA998565	c	
521	26119	AA998576	i,r,w,General	
522	22737	AA998660	aa	, , , , , , , , , , , , , , , , , , ,
523	3696	AA999030	е	
524	3079	AA999169	k,x,General	
525	3081	AA999171	e,p,r	
526	3082	AA999172	General	Glutamate metabolism, Purine metabolism
527	17337	AB000717	k	
528	1535	AB000778	а	
529	1382	AB002406	k	
530	20184	AB003753	d	
531	4312	AB010635	c,i,j,k,y,z	Mothiopino motabolism
532 533	21666	AB012214 AB015645	lk	Methionine metabolism
533 534	15772 1183	AF013144	g h	
535	1582	AF015144 AF015911	h,z	
536	11483	AF020618	u,cc,General	
537	20295	AF024712	aa	
538	19077	AF030358	y,z	
539	23044	AF034218	General	
540	25178	AF035955	d	
541	1564	AF035963	x,bb,General	
542	8426	AF036335	f	
543	21817	AF036537	k	
544	21145	AF038571	General	
545	22602	AF044574	General	
546	13464	AF047707	h	
547	24024	AF052695	X	
548 549	12259 4589	AF061266 AF062389	h y,z	
				1

vabus2: P	ZATIHIKWANYS	O A SPAN FOREST	34	ATTY: Docket No. 47921-5039W Doc. No. 1793397
Sequence ID No.	ldenfffer	ConBenk Ace/ Ref. Seq. ID No.	Model Gode	Pathways:
551	15761	AF062741	u	
552	17426	AF073839	р	
553	18615	AF074608	S	
554	15797	AF084205	f	
555	12932	AF102552	S	
556	18603	A1007649	х	
557	22733	AI007668	r	
558	22746	A1007672	r General	· · · · · · · · · · · · · · · · · · ·
559 560	24109 15848	AI007725 AI007820	n,v	
561	10108	Al007857	f	
562	6804	AI007877	General	-
563	20099	AI007893	f.u	
564	11368	AI007948	d	
565	15849	AI008074	h	
566	3121	AI008160	General	
567	16646	AI008190	t	
568	12683	AI008203	х	
569	22018	AI008309	b	
570	23917	AI008441	n Cananal	
571	22599	AI008458	General	
572	22698	A1008578	p,General	
573 574	14405	A1008579	r,x	
575	4086 3808	AI008629 AI008643	i.v.General	
576	3931	AI008697	I,v,Gerierai	
577	7785	Al008758	laa	
578	16701	AI008838	9	<u> </u>
579	21789	AI008930	k	
580	21895	AI008971	General	
581	410	AI008974	i,aa,General	
582	21632	AI009167	General	
583	21596	AI009168	General	
584	22801	AI009197	General	
585	11876	AI009321	cc,General	
586	2506	AI009341	General	
587	6382	AI009362	General	
588	14370	A1009427	k	
589	19275	A1009460	x	
590 591	4154 3464	AI009467 AI009589	g cc	
592	3926	AI009589	e	
593	19358	AI009592	c	
594	22545	AI009747	g	
595	15089	AI009752	cc.General	
596	5458	AI009756	h	
597	6844	AI009770	e,r,cc	
598	15627	Al009810	aa	
599	22619	AI009825	d	
600	7857	Al009898	j,l,m,z	
601	13259	AI009946	r	
602	21105	AI010067	General	
603	24627	AI010102	aa	
504	12716	AI010178	General	
505	18757	AI010216	aa	
506 207	2912	AI010220	aa,General	
607 608	3316 15644	AI010237 AI010256	t General	
609	657	AI010256	b	
610	3271	AI010202	b	
611	11081	AI010303	bb	
612	16521	AI010470	c,s,t,General	Porphyrin and chlorophyll metabolism
613	6927	AI010542	General	,

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TABLE 2: F				Aliy, Poetel No. 44921-5039W0 Doe, No. 1798397.1
ID No.	ldeniiier	Cenenk Ace Ref. Seq. ID No.	Model Gode	Pathways 1.4
614	17524	AI010568	a,j,y,General	
615	6946	AI010642	n	
616	23509	Al010962	aa	
617	6044	AI011285	t	
618 619	13855 21779	AI011361 AI011380	cc	
621	12534	AI011460	cc	
622	12629	AI011492	e.f	
623	735	AI011560	f	
624	3941	AI011598	General	
625	17550	AI011607	j,General	
626	10636	AI011634	e Canaral	
627 628	3995 16112	AI011678 AI011706	General h	
629	13354	AI011757	c	
630	12745	AI011799	cc	
631	18684	AI011812	t	
632	4205	Al011982	b	
633	6518	Al012114	General	
634	17407	AI012145	General	
635	13093	AI012177	r f	
636 637	15395 21796	Al012216 Al012221	d,General	10.0
638	3981	AI012235	i,General	
639	6606	AI012308	i,r	
640	3417	Al012337	w	
641	24200	AI012356	b,t,General	
642	7471	Al012379	cc	
643	7247	AI012438	9	
644	7127	AI012464	p,General	
645 646	2311	Al012471 Al012485	b aa	
647	20817	Al012589	g,n,q	Glutathione metabolism
648	3493	AI012590	v,General	
649	8975	AI012613	General	
650	11335	Al012619	j	
651	21409	Al012637	General	
652	8015	AI012638	aa	
653	8476 4232	AI012647	w e,p,General	
654 655	23128	Al012958 Al013011	General	
656	20086	Al013260	General	
657	11969	AI013273	k	
658	26147	Al013387	aa	
659	8815	Al013437	р	
660	19722	AI013508	k	
661	6674	AI013568	General	
662 663	23145 15130	Al013647 Al013676	o,t w	
664	7274	AI013715	aa	
665	7276	AI013713	e	
666	7278	AI013738	y,z,aa	
667	22592	AI013740	s,x,bb,General	
668	16584	AI013765	w	
669	24143	AI013804	j,l	
670	15928	AI013829	a,General	Valing lauging and igalousing degradation
671 672	21950 3260	AI013861 AI013875	t t	Valine, leucine and isoleucine degradation
673	2708	Al013882	d,q	
674	8585	Al013886	i	
675	7299	AI013911	p,r,t,General	
676	15904	AI013971	General	
677	12781	Al014023	w	

vasue2: F	ZASTIHAWASYS			Atity, Docket No. 44921-5089W Doc. No. 1798397
Sequence ID No.	ldentifier	Cenbank Aca Ref. Seq. ID No.	Model Gode	Pathways
678	19372	Al014135	aa	
679	4241	AI014140	w	
680	15247	AI014169	c,u	
681	7315	AI028831	n	
682	16631	AI028856	General	
683	23297	AI028953	X	
684	11326	AI029015	b	
685 686	2866 12812	AI029058 AI029126	n,y General	<u> </u>
687	17602	AI029126	p	
688	7392	AI029185	aa	
689	6517	AI029264	d,k,x	
690	7639	AI029292	b	
691	3874	AI029428	i,General	
692	12819	Al029437	f	
693	7452	Al029466	r	
694	7493	AI029608	b	
696	7537	AI029829	o,General	
697	2310	AI029969	V	
698	7585	AI030023	X	
699	7586	A1030024	b,n	
700 701	14492	AI030091	CC	
701 702	10673 7615	AI030134 AI030163	o,r	-
703	2370	AI030103	General	
704	7681	Al030449	n	
705	11559	AI030472	General	
706	7665	AI030668	t,bb	
707	24222	AI030704	k	
708	10740	Al030743	h	
709	10742	Al030773	е	
711	16169	AI030932	General	
712	19527	AI030991	f	
713	22614	AI031004	r	
714	3167	AI031012	e	
715 716	5350 7858	AI043611 AI043654	t t	
717	10784	AI043654 AI043678	d	
718	9180	AI043694	aa	
719	7867	AI043695	aa	Glutamate metabolism, Purine metabolism
720	7584	AI043724	General	C. C. C. C. C. C. C. C. C. C. C. C. C. C
721	7895	AI043768	e	
722	7903	AI043805	General	
723	7913	Al043849	cc	
724	3899	AI043904	1	
725	6766	AI043914	f	
726	10818	AI043990	g,I,m,General	
727	7956	AI044018	İ .	
728	5393	AI044170	р	
729	5398	AI044177	q	
730 731	5425 8692	AI044237 AI044247	a,d r	
732	5430	AI044247 AI044253	i -	
733	5461	AI044233	g.p.General	
734	5464	AI044336	i	
735	3359	AI044347	aa	
737	2695	AI044396	b	
738	5494	AI044425	General	
740	9882	AI044588	j,m	
741	5575	AI044688	g	
742	2348	AI044794	General	
743	18205	AI044836	n	
744	5626	AI044864	u	

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ABLE 2: [PATHWAYS		forgal and places, position	Air Docke: No. 44921-5039W0
			Production of the engineering	<u>Dos. No. 1798397.1</u>
D No. : Dequence	O-Convertions	Cenbenk Acel Ref. Seq. ID No.	Model Code	Pathways .
ບ ເນຍ _{ະເ} ຸ '45	Identifier 5630	A1044869	f	remmelys 3500
46	5634	AI044883	General	
47	4047	AI044947	l,m	
48	5654	AI044976	w	
'49	5684 19235	AI045056	General	
750 751	5689	AI045074 AI045075	i,aa,General	
752	5711	AI045151	General	
753	19237	AI045153	С	
754	9964	Al045161	f	
55	5735	AI045223	f	
756	5474 5811	AI045477	a,General	
'57 '58	5811	AI045502 AI045537	d,e General	
759	5839	AI045594	i	
60	6808	AI045600	s	
'61	17755	AI045608	у	
763	10020	AI045632	a	
64	5855	AI045669	General	
65 66	5881 5897	AI045789 AI045862	General	
67	5900	AI045866	y,z	
68	7540	AI045882	o,t,General	
769	5329	Al045970	р	
770	15093	AI058285	d	
771	8002	AI058304	_ <u> i</u>	
772	8017 6828	AI058341 AI058359	General	
773 774	8177	AI058539	aa	
775	3090	AI058730	aa	
76	10093	AI058746	g	
777	8143	AI058759	General	
778	18659	AI058762	f	
779	8163	AI058837	aa	
780 781	4789 8221	AI058889 AI059061	General General	
782	10159	AI059147	d	
783	8245	AI059154	b	
784	8283	AI059290	n	
785	8314	AI059386	g,General	
'86	10200	A1059444		
787 788	8347 18359	AI059519 AI059675	s n	
'89	10281	AI059947	b,t	
90	8494	AI059968	aa	
91	8495	AI059971	General	
92	8496	AI059974	General	
93	10289	A1060053	li la	
94	8548 8565	AI060176 AI060236	k t	
95 96	18322	AI060236 AI060279	i,y,z	
97	8745	AI069939	r r	
98	8785	AI070067	0	
99	17506	AI070068	cc	
300	9067	A1070087	General	
301	3551	A1070122	e	
302 303	4967 18	AI070179 AI070195	k General	
304	24197	AI070314	General	
305	8869	AI070330	r	
306	8874	A1070336	b,cc	
307	10417	Al070410		

TARILE OF I	TAVELUNAVIANX.C					
TABILE 2: P	ZAVIANWZANYS			Ality, Doeket No. 449241-50391110 Doe No. 1798397.1		
Sequence ID No.	legniffar	Cenbenk Ace Ref. Seq. ID No.	Model Gode	Pathways (*)		
809	14424	AI070421	I,p,General			
810	10434	AI070497	General			
811	8927	AI070523	V	· · · · · · · · · · · · · · · · · · ·		
812 813	8946 8950	AI070611 AI070621	q w			
814	8972	AI070673	General	 		
815	8981	AI070715	bb			
816	26184	AI070784	i,l			
817	3007	A1070824	w			
818 819	8999 10477	AI070839 AI070868	p e,f			
820	24301	Al0709011	k			
821	8721	AI071024	General			
822	9212	AI071098	x			
823	1831	AI071137	С			
824 825	11005 9104	Al071139	r			
826	9583	AI071173 AI071185	j,m General			
827	9644	Al071410	c			
828	16058	Al071490	General	Sphingoglycolipid metabolism		
829	11057	Al071509	f,o			
831	5695	AI071566	bb			
832 833	9671 22929	AI071568 AI071578	w General			
834	9673	Al071581	General			
835	9699	Al071646	General			
837	9799	Al072008	q,y,z			
838	9808	A1072050	d			
839	22796	Al072213	General			
840 841	9271 10869	AI072405 AI072425	w			
842	21797	Al072425	General			
843	9306	Al072521	r			
844	9312	Al072550	j			
845	10893	AI072559	x			
846	1501	AI072634	cc,General General			
847 848	9363	AI072658 AI072695	d			
850	9409	AI072841	n			
851	9410	AI072842	w			
852	9468	AI073021	General			
853	9518	AI073223	f	APA		
854	11183	AI100768	t	Nitrogen metabolism		
855 856	9190 2029	AI100835 AI100842	e p	 		
857	5687	Al101006	e			
858	15192	Al101099	g,cc			
859	17399	Al101157	0			
860	9339	AI101160	l,m,o			
861 862	6321 5421	Al101256 Al101270	General			
863	11910	Al101270 Al101323	General			
864	23140	AI101608	е			
865	4119	Al101901	General			
866	16324	Al102009	b			
867	18642	AI102023	0			
868 869	19373 7051	Al102044 Al102055	h			
870	6544	AI102064	C			
871	10227	AI102248	w			
872	23849	Al102318	e,q			
873	11954	AI102505	g,j,s	Oxidative phosphorylation		
874	2125	AI102519	c,k			

TABUE 2: PATHWAYS Doc No. 179					
Ecquence	ldentifer	Geneank Acc/ Ref. Seq. ID No. #	Model Code	Pathways	
875	5967	Al102520	У	- constitute and the Aller	
875	5969	Al102520	p,w		
876	11563	Al102560	General		
877	15190	Al102562	b,g,n,p,v		
878	19769	Al102570	bb		
879	22487	Al102578	General		
880	19011	A1102618	General		
881 882	23837	AI102620	q,t g,General		
883	23538 17234	AI102727 AI102741	c c		
884	5891	Al102741	k		
885	6796	AI102753	General		
886	8837	Al102849	0,p		
887	15861	Al102868	i		
888	3533	AI102877	g		
889	13222	AI102977	General		
890	6806	AI103018	o,u		
891	10659	Al103059	w,cc,General		
892	17400	Al103097	е		
893	3584	AI103106	x,aa		
894	13298	Al103143	r		
895	15981	Al103150	i,x		
896	3475	Al103245	w		
898	23619	AI103314	р		
899	24181	Al103320	e		
901	4355	AI103410	General General		
902 903	7622 20918	AI103472 AI103552	n		
904	21579	AI103532 AI103572	General		
905	2222	AI103572 AI103631	0		
906	2752	AI103641	e		
907	4856	AI103708	li .		
908	8990	Al103719	l,m,y,z		
909	15942	AI103738	г		
910	22885	AI103828	e,General		
911	15853	AI103841	x		
912	15050	Al103911	j,y	Oxidative phosphorylation	
913	12376	Al103939	U		
914	22271	Al103947	o,y		
915	20833	Al104035	f,q	Oxidative phosphorylation	
916	7010	Al104099	w		
917	22101	AI104251	General		
918	22833	AI104258	General		
919 920	22211 10720	AI104279 AI104296	g,m		
920	15416	A1104296 A1104340	<u> </u>		
922	10991	AI104340 AI104342	a		
923	18831	AI104342 AI104357	p		
924	7223	AI104377	e		
925	23574	Al104520	e,g,s	Oxidative phosphorylation	
926	18509	Al104528	q	2500 15 000 000 000 000 000 000 000 000 0	
927	11680	AI104605	v v		
928	12342	AI104658	w		
929	23689	AI104685	r		
930	15377	Al104821	o,cc		
931	22957	Al104897	General	Oxidative phosphorylation, Type III protein	
932	18451	AI104953	o,s	secretion system	
933	24375	Al104979	n,General		
934	18278	Al105080	bb		
935	2196	Al105243	g		
936	5199	Al105272	bb,General		
937	12901	Al105301	o,s		

TABUE 28 F	PATHIKKAYYS	3		A ti y, Docket No. 44921-5039W.c Doc: No. 1798397.
Ecquence ID No.	ldeniiier	Cenbenk Ace Ref. Seq. ID No.	Model Gode	Pathways 4
938	7700	AI105383	cc,General	
939	13343	AI105398	u	
940	22931	AI105417	e,General	
				Fatty acid metabolism,Lysine
941	23596	AI105435	bb	degradation,Tryptophan metabolism
942	15893	Al105465	0	
943	12660	Al111492	C General	
944 945	4479 24211	Al111599 Al111853	k	
946	2539	Al111960	r	
947	5729	AI111990	k	
948	4049	Al112012	i,g.u,General	
949	12908	AI112043	i	
950	20041	Al112161	t	
951	12937	Al112462	General	
952	3713	Al112571	b	
953	12921	Al112636	General	
954	12965	Al112926	General	
955	7499	Al112986	General	
956	4969	AI113008	r	
957	11817	AI136295	f	
959 960	11165 4045	AI136372 AI136460	cc	
961	12782	Al136493	k	
962	6850	Al136665	h	Purine metabolism, Pyrimidine metabolism
963	20920	Al136891	p,v	Turne metabolism, yilmane metabolism
964	6552	Al137062	0	
965	22722	Al137211	i	
966	13111	Al137224	o,General	
967	15969	Al137302	е	
968	14349	Al137303	d	
969	9166	Al137406	General	
970	9525	Al137516	r	
971	6638	Al137579	General	
972	7414	AI137586	General	
973 974	11321	AI137752	Z	<u> </u>
974 975	23473 13158	AI137932 AI138024		
976	13467	Al138034	cc	
977	11377	Al138105	V	
978	6790	Al144801	d,h	
979	6506	Al144919	j,l,y	
980	8027	AI144958	i i	
982	14458	Al145095	General	
983	7476	Al145202	g	
984	17545	AI145384	е	
985	17479	AI145385	Γ	
986	4194	Al145387	r	
987	8634	Al145722	g	
988	8339	Al145761	y,General	
989	2059	AI146005	h,General_	
990	23224	AI146033	O bb	Valina lausing and indeputing described
991 992	5232 18472	AI168942 AI168975	bbu	Valine, leucine and isoleucine degradation
992	18473	AI168975	u	
993	13235	AI169020	r	
994	11618	AI169115	o,y,General	
995	17386	Al169144	0	
996	10984	AI169156	o,u	
997	8205	Al169176	е	
998	12979	Al169177	е	
999	2607	Al169211	С	

û∖van∃ sæ l	ATHWAYS			/- Xiiy, Docket No. 44921-5039 Doc. No. 179339		
Sequence ID No.	ldeniffer	Cenbank Acel Ref. Seq. ID No.	Model God:	Pathways		
1000	22661	AI169265	6.7	Oxidative phosphorylation, Type III protein secretion system		
1001	13239	Al169278	g,j,l,y,z	secretion system		
1002	24162	AI169279	m	- 		
1003	16879	Al169284	0			
1004	24213	Al169289	р			
1005	13240	AI169311	СС			
1006	5931	Al169324	b			
1007	20891	Al169337	d			
1008	11979	AI169365	cc			
1009 1010	10947	AI169372	s			
1011	20697 8234	AI169494 AI169517	o,u z	******		
1012	18343	AI169648	0			
1013	10839	Al169655	I,m			
1014	24146	Al169668	j,l			
1015	22575	Al169728	r			
1016	804	Al169756	cc			
1017	8213	Al169883	р			
1018	3916	Al169947	i,bb			
1019	3733	Al170053	u,General			
1020	14179	AI170224	cc			
1021	11406	Al170263	r			
1022	3547	Al170279	General			
023	11524	AI170340	j,y,z			
024	2729	AI170363	e,i			
1025	18811	AI170525				
1026 1027	22524 24048	Al170542 Al170570	h			
1028	5968	AI170570	a,g y,aa			
1029	9757	AI170693	b			
1030	18905	AI170770	e,s			
1031	16170	AI170894	i			
1032	7089	AI171185	c			
1033	17591	AI171354	b			
1034	13285	AI171361	h			
1035	4428	Al171362	а	Oxidative phosphorylation, Ubiquinone biosynthesis		
1036	18126	Al171369	w			
037	23253	Al171448	0			
1038	4584	AI171492	m,General			
1039	11158	AI171542	r,s			
1040	15345	AI171587	<u> </u>			
1041	21183	AI171676	k			
1042	8215	AI171692	i			
043	11437 2625	Al171794 Al171800	i			
045	23579		cc			
046	11708	Al171802 Al171807	l,t			
047	17204	AI171844		Oxidative phosphorylation,Type III protein secretion system		
047	4420	AI171916	s,y,z m	Scoretion system		
049	3266	Al171948	I,m			
050	19012	AI172056	t			
051	11205	Al172057	a,q,bb			
052	6057	Al172102	b			
053	19128	Al172103	m			
054	15673	Al172107	z			
055	6630	AI172184	n			
056	11968	AI172208	bb			
057	6974	AI172263	l,m			
058	23313	AI172271	d			
059	2140	AI172272	General			

TVABLE 2: P	AYAWKHTKA			/ Aliy. Docket No. 44921-5089//0 Doc. No. 1798397.1
Sequence ID No.	ldeniiier	ConBank Acel Ref. Seq. ID No.	Model Gode	Pathways
1060	15382	Al172302	l,p,General	
1061	18689	Al172329	l	
1062	17887	Al172414	0	
1063	3042	AI172447	General	O'L to a de (TOA a and a) O'L to the land
				Citrate cycle (TCA cycle), Glutathione metabolism, Reductive carboxylate cycle
1064	17291	AI172491	bb	(CO2 fixation)
1065 1066	26222 13095	AI172506 AI172595	r	
1067	8795	AI172618	General	
1068	6454	Al175342	j,l,m,y	
1070	4445	AI175466	х	
1071	3418	Al175475	m	
1072	18507	Al175551	bb	
1073	10217	Al175628	w	
1074	7262	AI175833	j,m,x	
1075	19004	AI175875	I General	
1076 1077	22352 7022	AI175959 AI176041	l,General h,n	-
1077	21467	AI176041	111,111 t	
1078	18581	AI176160	General	
1080	14159	AI176169	g	
1081	21742	AI176172	w	
1082	10182	AI176185	V	
1083	22765	Al176265	General	
1084	6905	Al176275	а	
1085	12999	AI176276	cc	
1086	16438	AI176294	е	
1087	21130	A1176298	У	
1088	3014 15015	AI176362 AI176363	e	
1089 1090	19006	Al176393	x	
1090	20001	A1176396	6	
1092	12174	AI176435	i,m	
1093	15191	Al176456	b,o,t,v,cc	
1094	24236	Al176473	d,General	
1095	16518	AI176546	v	
1096	2161	AI176592	General	
1097	12436	AI176610	General	
1098	2536	AI176616	I,v,General	
1099	18525	AI176792 AI176828	u	
1100	23449	AI176839	g General	
1101 1102	3580	Al176848	e	
1103	22103	AI176849	d.General	
1104	16036	AI176855	f	
1105	15588	Al176916	General	
1106	16917	Al176951	t	
1107	16124	AI176963	СС	
1108	15146	Al176969	b,General	
1109	5786	AI177058	f	
1110	2852	AI177059 AI177092	С	
1112 1113	3156 14384	Al177092 Al177096	g a	Purine metabolism
1114	13310	AI177119	General	T dillo metabolism
1115	24049	AI177341	g,p,s,u	<u> </u>
1116	15964	AI177360	o,General	
1117	14989	AI177366	u	
1118	7975	Al177374	aa	
1119	3006	AI177395	k	
1120	17570	AI177683	r	
1121	9521	AI177706	b	
1122	14425	AI177755	g,General	

TABUE2: P	SYAWHTA	\$ 1.36		Afiy. Docket No. 44921-5039W Doc. No. 1793397
Sequence ID No.	ldeniiier	GenBenk Acci Ref. Seq. ID No. "	Model Gode	Pathways
1123	10611	AI177790	j,m	
1124	5356	Al177813	cc	
1125	11791	Al177843	General	
1126	14484	AI177867	General	
1127 1128	5780 19184	Al177869 Al178025	General General	
1129	6059	AI178025	c,General	
1130	23248	Al178267	v	
1131	4073	Al178272	o	
1132	7838	Al178291	е	
1133	18996	Al178326	у	
1134	22488	Al178392	b	
1135	18800	AI178504	n,p,aa	
1136	22197	AI178527	g,General bb	
1137 1138	3401 17713	Al178684 Al178700	m	
1139	14874	AI178735	e	
1140	23567	AI178746	v,General	
1141	18907	AI178971	С	
1142	20991	Al178979	i	
1143	5887	Al179099	q,t	
1144	8477	Al179167	b,e,General	
1145	3348	AI179288	u,v	
1146	13608	AI179314	е	
1147	8849	AI179315	g,p v,General	
1148 1149	13611 15438	AI179378 AI179399	m,x	
1150	13614	AI179407	e,t,General	
1151	15042	Al179422	b,General	
1152	2768	Al179481	i,General	
1153	24041	Al179580	b,i	
1154	19822	Al179599	o,General	
1155	23270	AI179601	q,General	
1156	5901	AI179605	e	Porphyrin and chlorophyll metabolism
1157 1158	16081 14564	AI179610 AI179717	g,i,p k	Porphyrin and Chlorophyli metabolism
1159	7918	AI179750	General	
1160	6647	AI179795	g	
1161	9097	Al179875	o,General	
1162	23989	AI179953	а	
1163	12899	Al179967	b	
1164	1687	Al179971	С	
1165	22569	AI179979	General	Chaire series and there is a series are to be a
1166	23514 15892	AI179986 AI179988	o,General c,General	Glycine, serine and threonine metabolism
1167 1168	12402	AI180004	g	
1169	5443	AI180165	General	·
1170	5481	AI180170	General	
1171	24028	Al180239	1	
1172	17089	AI180281	g	
1173	3701	Al180306	aa	
1174	3352	AI180334	m	
1175	24368	Al180392	l,m	
1176 1177	14337 19080	Al180414 Al227647	C LV7	
1178	22838	Al227647	j,y,z aa	
1179	6765	Al227761	i,General	
1180	24054	Al227867	General	
1181	7324	Al227885	i	
1182	23898	Al227987	d	
1183	1651	AI228068	n,w	
1184	14237	AI228128	e	
1185	14242	Al228197	General	i

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TABUE 2: P	AYAWHIVA		51 miles	/X ii y, Docket No. 44921-5089WG Doc. No. 1798397,
Sequence ID No.	ldenillier	ConBank Acc Ref. Seq. ID No.	Model Gode	Pathways : 1
1186	16913	AI228236	0	
1187	22915	AI228299	r Canada	
1188 1189	8917 15879	AI228301 AI228313	General r,General	
1190	13727	Al228326	o,General	
1191	6102	AI228335	General	
1192	13730	AI228356	а	
1193	13745	AI228494	b,cc	
1194	4217	Al228587	s	
1195	16053	AI228596	СС	
1196 1197	3557 11605	Al228672 Al228682	e e	
1198	13203	AI228728	r	
1199	13771	AI228848	g	
1200	5918	AI229036	r	
1201	8235	Al229154	k	
1202	16203	Al229196	r	
1203	13826	AI229304	a	
1204 1205	13144 4640	Al229320 Al229404	g	
1206	23563	Al229404 Al229421	x,aa	
1207	15426	Al229421	s	· · · · · · · · · · · · · · · · · · ·
1208	15193	Al229508	bb	
1209	19243	AI229638	х	
1210	23078	Al229647	р	
			\	Oxidative phosphorylation, Ubiquinone
1211	3099	AI229680	0	biosynthesis
1212 1213	19508 13977	Al229698 Al229707	bbx	
1213	23983	AI229707 AI229708	V	
1215	2688	Al229793	e	
1216	13874	Al229832	g	
1217	12587	Al229979	General	
1218	20591	AI229993	l,m	
1219	24042	AI230002	a,b,d,General	
1220	13880	AI230042	u	Oxidative phosphorylation, Ubiquinone
1221	17672	AI230074	d	biosynthesis
1222	3652	Al230113	General	biosynthesis
1223	18650	Al230121	aa	
1224	13025	Al230173	С	
1225	4280	AI230247	z	
1226	18528	Al230284	General	
1227	7084	AI230362	p	
1228 1229	20895 12961	Al230549 Al230554	b,n General	
1230	15636	Al230504 Al230616	r	-
1231	4121	Al230647	j,m	
1232	14388	Al230702	General	
1233	18529	Al230716	x,General	
1234	13618	Al230724	General	
1235	8304	AI230746	cc	
1236	4731	A1230773	e	
1237 1238	14430 16627	Al230798 Al230822	c,k,x bb	Glycoprotein biosynthesis
1239	3125	AI231028	General	City oop rote in biodynthesis
1240	633	AI231020	k	
1241	20846	Al231140	p	
1242	6743	AI231219	d	
1244	26292	Al231391	k	
1245	12343	AI231433	w	
1246	7337	AI231465	General	
1247	16321	AI231506	General	J

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TABLE 2: F				/4447. Doorot No. 44921-5089W. Dog. No. 1798997.
Sequence ID No. :	ldeniller	CenBenk Acel Ref. Seq. ID No.	Model Gode	Pathways *
1248	8004	Al231532	j,l	
1249	15171	Al231792	g	
1250	6193	Al231797	ļi	
1252	14227	AI231999	u	
1253	24501	AI232006	w,y,bb	
1254	3434	AI232014	g,q,z,cc, General	
1255	19094	AI232021	n,General	
1256	14020	AI232076	u	
1257	6726 11549	AI232157 AI232174	l,m	
1258 1259	23125	Al232174 Al232266	j,s	
1260	2085	Al232270	bb	
1261	2913	Al232270 Al232272	0	
1262	14304	Al232281	g	
1263	15955	AI232294	u,bb,General	
1264	15122	AI232303	y	
1265	4716	AI232303	y	
1266	15246	AI232332	t,u	
1267	24321	AI232340	0	
1268	16172	AI232341	d	
1269	11411	AI232346	h	
1270	19287	Al232379	f	pdgf
1271	5601	AI232461	n,General	1
1272	14051	AI232489	l,m	
1273	5572	Al232490	i,t	
1274	11157	Al232494	cc	
1275	8709	AI232534	0	
1276	20350	AI232552	j,v,y	
1277	14069	Al232631	е	
1278	4440	Al232643	w	
1279	17695	Al232784	е	
1280	15796	Al232874	v	
1281	12467	Al232924	General	
1282	12873	Al232984	i	
1283	5355	Al233031	r	
1284	18794	Al233121	С	
1285	3823	AI233147	b,g,General	
1286	11967	AI233155	c,k,General	
1287	11561	Al233182	d	
1288	3471	AI233183	9	
1289	21948	AI233191	0.01/	
1290	13598	AI233194	g,p,y	
1291 1292	15552 17907	Al233195 Al233224	bb	
1293	14111	AI233224 AI233269	CC	
1293	12894	AI233269 AI233365	d	
1295	7161	AI233407	General	
1296	15906	AI233407 AI233425		
1297	14120	AI233425	d d	
1298	14095	AI233458	a,d	
1299	3075	Al233494	u,aa	
1300	6046	Al233530	General	
1301	18900	Al233570	General	
	1			Aminoacyl-tRNA biosynthesis, Arginine and
1302	7888	AI233583	General	proline metabolism
1303	16709	AI233602	General	Purine metabolism
1304	5163	AI233712	у	
1305	7243	AI233717	General	
1306	3816	AI233729	9	
1307	13023	AI233740	d,h,General	
1308	14871	AI233743	g	
1309	7469	AI233767	cc	

ITABLE 2: PATHIWAYS					
	ZAYZANWANYS			Affy: Docket No. 44924-5039VV(Doc. No. 1798397,	
Sequence ID No.	ldemilier	Cenbank Ace Ref. Seq. ID No.	Model Gode	Pathways 4	
1310	7804	Al233771	b		
1311	13563	AI233773	е		
1312	2154	AI233818	k,cc		
1313 1314	16616 13393	Al234079 Al234100	h a,d,General		
1315	7071	Al234162	r		
1316	14677	AI234620	General		
1317	4443	AI234629	m		
1318	22453	AI234678	b		
1319	23964	AI234748	t,General		
1320 1321	19581 22152	Al234753 Al234822	o,General		
1322	18942	AI234865	d		
	,,,,,			Oxidative phosphorylation, Type III protein	
1323	22662	Al234939	aa	secretion system	
1324	3875	Al235047	o,General		
1325	19479	AI235135	0		
1326 1327	14906	AI235192	g		
1327	14718 15004	Al235210 Al235224	e b,General		
1329	6632	Al235277	v		
1330	14722	Al235284	x,z		
1331	1462	Al235585	u,General		
1332	21061	Al235631	l,m		
1333	14665	Al235646	m	tgf-beta	
1334	19940	AI235689	General		
1335 1336	5698 23745	Al235692 Al235732	u k		
1337	11164	Al235732 Al235739	General		
1338	5212	Al235745	d		
1339	14768	Al235912	h		
1340	14776	AI235950	m		
1341	3091	AI236027	n,General		
1342	14861	AI236045	r		
1343 1344	14862 16943	AI236048 AI236097	e p		
1345	8336	Al236101	1		
1346	23230	Al236146	v		
1347	22855	Al236150	е		
1348	14594	Al236152	i		
1349	18406	AI236168	r		
1350	15051	A1236332	General		
1351 1352	19298 10667	AI236338 AI236366	bb b		
1353	10774	AI236397	f		
1354	9407	AI236402	aa		
1355	26335	AI236460	General		
1356	17950	Al236590	t,General		
1357	18259	AI236601	h,v		
1358	11445	AI236613	j,y		
1359 1360	17248 16859	AI236635 AI236753	o,aa t,General		
1361	5208	Al236754	g		
1362	24388	Al236772	e,General		
1363	15850	Al236795	n,v,w		
1364	14800	Al236856	w		
1366	11404	AI237002	m		
1367	18151	AI237212	o,General		
1368 1369	21653 11208	Al237535 Al237586	t,General z		
1370	21893	Al237713	i,k,aa		
1371	14842	Al237724	r		
1372	3467	Al237835	General		

TABLE & P	ATHWAYS			/Xiiy, Dooket No. 4/1924 -5039/// Doc. No. 1798397.
Sequence ID No.	ldentifer	Coneank Acal Ref. Soq. ID No.	Model Gode	Pathways to the state of the st
1373	25840	Al638972	u	
1374	17108	AI639017	ń	
1375	16676	A1639082	c,k,x	
1376	12400	AI639107	k	
1377	19952	AI639108	q,v	
1379	25907	AI639167	o,w	
1381 1382	18533 18353	AI639231 AI639233	n t,aa	
1384	15330	Al639285	General	
1385	20026	AI639354	g	
1386	25971	AI639365	r	
1388	19152	AI639387	u,General	
1390	18338	AI639422	y	
1392	20082	AI639488	i,m	
1394	20056	AI639504	a,bb,General	
1395	4713	AI639518	q	
1396	14332	AJ001044	bb	
1397	7602	AJ001929	k	
1398	9867	AJ005424	u	999
1400	16351	AJ011811	General	
1401	20116	AJ011969	I,General	
1402 1403	17635 18686	AJ223355 D00729	v,w	Fatty acid metabolism
1404	5049	D10655	n,w	Fatty acid metabolism
1405	25257	D13623	i	*
1405	15281	D13623	h	
1406	11434	D14014	cc	
1407	1613	D14076	x	
1408	1728	D16479		Bile acid biosynthesis, Fatty acid biosynthesis (path 2), Fatty acid metabolism, Phenylalanine metabolism, Valine, leucine and isoleucine degradation
1409	3015	D16554	c,s,v,z	degradation
1410	472	D26111	d,s,bb	
1412	16233	D29960	i.l	
1413	9029	D30804	n	
1414	1485	D38222	y,z	
1415	9135	D45247	s	Proteasome
1416	16354	D50564	u	Cysteine metabolism
1417	1884	D50695	l,m,bb	
1418	21147	D63772	General	
1419	826	D82928	ĮŤ	Glycerolipid metabolism
1420	25306	D84485	u	
1421	18867	D88250	t Conoral	
1423 1424	22543 12360	H31117 H31456	r,v,General w	
1424	20514	H31489	h,j	
1426	11358	H31610	h	
1427	4360	H31813	bb,General	
1428	9343	H32169	1	-
1429	4386	H33093	h,w	
1430	4415	H33636	h	
1431	15374	H34186	l .	· · · · · · · · · · · · · · · · · · ·
1432	17159	J00797	u,General	
1433	16260	J01878	f	
1434	17284	J02827	bb	Valine, leucine and isoleucine degradation
1435	15017	J03752	n	
1436	44	J03819	p,s	
1437	21014	J03914	e,r,General	Glutathione metabolism
1438	20429	J05035	f	Androgen and estrogen metabolism,Bile acid biosynthesis

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vabue2: v	PATHIWAYS			ATTY, Doctot No. 449214-5039 Doc. No. 17983
Sequence ID No.		Cenbenk Acel Rel Seg. D No.	Model Code	Palliways
1439	1247	J05181	j,l,m,s,y,z	Glutamate metabolism, Glutathione metabolism
1440	10464	J05510	n,u,General	
1441	20149	K03243	9	
1442	17758	K03249	9	
1443 1444	381 2048	L00124 L00382	k,x	
1445	10500	L04619	s	<u> </u>
1447	108	L14002	р	
1448	25366	L14003	t	
1449	109	L14004	c,p	
1450 1451	20414 25369	L14323 L14937	General	
1452	16119	L16532	k	
1453	25377	L25387	h	
1453	12058	L25387	h	
1455	21146	L35558	General	
1456	106	L37203	w	
1458	13682	L38482	f,j,k,m,z	Glutamate metabolism, Glutathione
1459	6405	L38615	p	metabolism
1461	15189	M11794	n,v	THO CLOUIS THE
1462	17086	M13011	j	
1464	21053	M15481	0	
1465	25405	M18330	j,l	
1466	25415	M19648	а	
1468	14967	M22366	bb	
1469 1471	20481 15048	M22631 M24542	q	Oxidative phosphorylation
1472	20921	M29853	m	Caldative prioapriorylation
1473	1224	M31931	u	
1474	15579	M33648	q	
1474	15580	M33648	q	
1475	17211	M34331	g,n,q,v	
1476	20699	M35601 M35601	b,x,bb b,t,bb	
1476 1477	9223	M36151	0	
1479	1585	M57728	j,m,y	
1480	24844	M58040	c	
1481	25057	M58495	h	
1482	457	M60666	d,General	
1483	1223	M75281	i k oo	ļ
1484 1485	5733 4198	M81855 M83143	i,k,aa m	
1485	4199	M83143	m	
1486	24651	M83678	k,x,z	
		,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,		Histidine metabolism,Phenylalanine metabolism,Tryptophan
1487	1430	M84648	General	metabolism, Tyrosine metabolism
1488	25467	M93297	С	Arginine and proline metabolism, Urea c and metabolism of amino groups
1489	729	M95762	a,y	
1490	23698	NM_012489	q	Bile acid biosynthesis, Fatty acid biosynthesis (path 2), Fatty acid metabolism, Phenylalanine metabolism, Valine, leucine and isoleuci degradation
1490	23699	NM_012489	q	Bile acid biosynthesis,Fatty acid biosynthesis (path 2),Fatty acid metabolism,Phenylalanine metabolism,Valine, leucine and isoleuci degradation

IJABLE 28	PATHWAYS			Atiy, Docket No. 44921-5039W Doc. No. 1793397
Seguence		ConBank Acc		Edge Wet 1// Sessol
D No.	refillmebl	Ref. Seq. ID No. *		Pathways
				Carbon fixation, Fructose and mannose
				metabolism,Glycolysis /
				Gluconeogenesis,Inositol
1491	7062	NM_012495	<u>q</u>	metabolism,Pentose phosphate cycle Fructose and mannose
				metabolism,Galactose
				metabolism, Glycerolipid
				metabolism,Pentose and glucuronate
1492	15511	NM_012498	u	interconversions,Pyruvate metabolism
1494	7427	NM_012515	General	
1495	24433	NM_012527	 	Arginine and proline metabolism,Urea cycl
1496	4467	NM 012529	d	and metabolism of amino groups
1497	16520	NM 012532	General	Porphyrin and chlorophyll metabolism
1498	225	NM_012544	x,z	
·				Histidine metabolism,Phenylalanine
			<u> </u>	metabolism,Tryptophan
1499	1431	NM_012545	General	metabolism,Tyrosine metabolism
1500 1500	23868 23872	NM_012551 NM_012551	I,m,v,General	
1500	23869	NM_012551	v.General	
1000	20000	1411_012351	1,00noral	Glycolysis /
		_A_A		Gluconeogenesis, Phenylalanine, tyrosine
1501	19407	NM_012554	z	and tryptophan biosynthesis
				Glycolysis /
1504	10100	NINA 040554		Gluconeogenesis, Phenylalanine, tyrosine
1501 1502	19408 21836	NM_012554 NM_012555	n,s,y,z k	and tryptophan biosynthesis
1502	21000	14141_012555	N	Carbon fixation,Fructose and mannose
		1		metabolism, Glycolysis /
1503	16895	NM_012558	g,s	Gluconeogenesis,Pentose phosphate cycl
1504	25317	NM_012559	bb	
1504_	6477	NM_012559 NM_012559	b,bb bb	
1504 1505	6478 11731	NM_012561	k	
1507	4254	NM 012564	a	
1508	16026	NM_012578	r	
1508	16024	NM_012578	r	
1508	16025	NM_012578	r	
1509	16080	NM_012580	g,m	Porphyrin and chlorophyll metabolism
1510 1511	15098 4450	NM_012588 NM_012592	bb	Valine, leucine and isoleucine degradation
1511	4451	NM_012592	i,bb	Valine, leucine and isoleucine degradation
1511	4452	NM 012592	bb	Valine, leucine and isoleucine degradation
1512	17198	NM_012593	a,x	
512	17197	NM_012593	х	
1513	18749	NM_012600	a,h	Carbon fixation,Pyruvate metabolism
1514 1514	2628 2629	NM_012603 NM_012603	General x,General	
1515	16849	NM 012608	n,o,q	
1517	15540	NM_012620	General	
1518	24568	NM_012630	General	
518	24566	NM_012630	General	
519	18553	NM_012631	k General	
1520 1521	1844 24668	NM_012637 NM_012642	General f	
1522	18632	NM 012645	a	
1523	25435	NM_012647	g	
1524	9423	NM_012649	b,cc	
525	24496	NM_012654	n	
526	7101	NM 012679	x,bb,General	
1320		11111_012010		Fatty acid metabolism, Tryptophan

VABUE 2	PATHWAYS			ATTY, DOGRETINO, 44924-5039WC
				<u>Dog. No. 1793397.1</u>
Sequence D No.	9	Genbenk Acc Ref. Seq. ID No.	Model Code	(Fothways)
			Money Cone	Peunways
1528 1528	1850 1854	NM_012696 NM_012696	t	-
1529	1603	NM 012697	General	
_				Aminosugars metabolism, Fructose and mannose metabolism, Galactose metabolism, Glycolysis / Gluconeogenesis, Starch and sucrose
1530	1372	NM_012734	u	metabolism
1531	1478	NM 012744	bb,General	Alanine and aspartate metabolism, Citrate cycle (TCA cycle), Pyruvate metabolism
1532	343	NM 012747	h,t	Cycle (10/10/cle),i yravate metabolism
1533	8829	NM_012749	General	
1534	20828	NM_012752	General	
1534	20829	NM_012752	i,General	
1534	20830	NM_012752	i,General	
1535 1536	15174 21685	NM_012756 NM_012760	j,m,n	
1537	18068	NM_012762	t	
1538	1246	NM 012770	a,General	Purine metabolism
1539	1348	NM_012776	f	
1540	18135	NM_012791	w	
1541	16947	NM 012793	p,bb	Arginine and proline metabolism, Glycine, serine and threonine metabolism, Urea cycle and metabolism of amino groups
1542	960	NM 012796	u	Glutathione metabolism
1543	260	NM_012798	f,u	
1544	556	NM_012803	d	
1545	21729	NM_012804	q	
1546	15032	NM_012816	General	
1547 1548	24895 18109	NM_012817 NM_012823	General u,General	
1549	373	NM 012833	h,l,q,General	
1550	2855	NM 012838	е	
1551	11136	NM_012839	S	
1552	20885	NM_012842	а	egf
1552	20884	NM_012842	a,bb	egf
1553 1554	18770 20674	NM_012857 NM_012861	e	
1554 1555	13151	NM 012862	a,r,General	
1556	24617	NM 012870	General	
1557	20945	NM_012875	a,v	
1558	15872	NM_012879	o,r	
1559	495	NM_012880	Z	
1559	494 23651	NM_012880	d u Conoral	
1560 1562	19477	NM_012881 NM_012891	d,u,General q	
1563	18564	NM_012899	v,General	Porphyrin and chlorophyll metabolism
1564	7197	NM_012904	f,r,cc,General	
1564	7196	NM_012904	v,cc,General	
1565	20202	NM_012909	b,r	
1566 1566	16581	NM_012911	c,j	
1566 1567	16582 24431	NM_012911 NM_012912	c General	
1568	18118	NM 012913	p	Oxidative phosphorylation
1569	6108	NM_012915	n	
1570	20757	NM_012923	c,i,aa	
1570	20755	NM_012923	i	
1571	2830	NM_012925	ļţ.	
1571 1572	1977	NM_012925 NM_012930	q	Fatty acid metabolism, Glycerolipid metabolism
1573	18694	NM_012931	j,l,m,z	
1574	13723	NM 012935	n	

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TABUE 2: F	ATHWAYS			Atty, Docket No. 42921-5089VV Doc. No. 1798997.
Sequence : ID No. : : :	ldentifer	CenBank Acel Ref. Seq. ID No.	Model Gode	Pailwelys
1575	9109	NM_012939	j,y,z	
1575	19398	NM_012939	aa	
1576	223	NM_012945	b,cc	
1577 1579	15058	NM_012950 NM_012963	cc	
1579	19374	NM 012964	g x	
1581	2554	NM 012967	t	
1581	2555	NM 012967	t,cc,General	
1582	24528	NM_012973	С	
1583	956	NM_012976	С	
1584	16417	NM_012991	g	
1585	17393	NM_012992	d	
1586 1587	23544 1588	NM_013013 NM_013026	s k	
1588	17894	NM 013026	m	
1589	18300	NM 013030	s,v,General	
1589	18076	NM 013030	g,s,z	
1589	18078	NM_013030	s	
1589	18077	NM_013030	e,s,z	
1591	730	NM_013040	w	
1592	17401	NM_013043	i,o,General	
1593	16684	NM_013052	General	
1594	14421	NM_013053 NM_013058	k	
1595 1596	15254 14997	NM 013059	s,z	Folate biosynthesis, Glycerolipid metabolism
1596	14996	NM 013059	General	Folate biosynthesis, Glycerolipid metabolism
1597	25676	NM 013069	aa	
1597	16924	NM_013069	0	
1598	24748	NM_013070	h,q	
1599	1529	NM_013082	d,General	
1600	1521	NM_013091	j,l,z,General	
1601	1685 26150	NM_013096 NM_013096	c,aa	
1601 1601	1688	NM 013096	c,i.	
1601	1689	NM 013096	c,p	
1601	1684	NM 013096	c,s,aa	
1602	20886	NM_013097	u,x,bb	
1602	20887	NM_013097	u,x,bb	
1603	1321	NM 013098	c	Galactose metabolism, Glycolysis / Gluconeogenesis, Starch and sucrose metabolism
1604	15296	NM_013102	l,m	
1606	23709	NM_013113	o,s,z,aa	
1606	23711	NM_013113	р	
1606	23710	NM_013113	s	
1607	1976	NM_013118	u	
1609	870	NM_013130	h Conoral	
1610 1611	16650	NM_013132 NM_013134	u,General	Storal hipsynthesia
1611	650 651	NM_013134 NM_013134	h h,j,l	Sterol biosynthesis Sterol biosynthesis
1612	1712	NM_013138	General	otoror biodyntalesis
1613	16982	NM_013144	o,v,General	
1614	21683	NM_013154	t,cc,General	
1614	21682	NM_013154	cc	
1615	3431	NM_013156	b,g,n	
1615	25567	NM_013156	v,General	
1615	3430	NM_013156	General	
1616	1309	NM_013159	w	
1616 1617	1310 21723	NM_013159 NM_013174	w w	
1618	1314	NM_013174 NM_013181	m	
1619	17357	NM 013183	p,bb,General	

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TABUE 2: (PATHWAYS			Aliy. Dockt No. 44921-508900 Doc. No. 1798397.1
Sequence ID No.	(Identiffer	Conbank Acel Ref. Seq. ID No.	Model Gode	Pathways VIII (VIII)
				Fructose and mannose metabolism,Galactose metabolism,Glycolysis /
1620	1300	NM_013190	у	Gluconeogenesis, Pentose phosphate cycle
1621	16448	NM_013197	С	Glycine, serine and threonine metabolism
1622	20856	NM_013200	b	Fatty acid metabolism, Glycerolipid metabolism
1623	397	NM_013214	f	THO COOKEN
1624	20864	NM_013215	g,n,y	
1625 ·	20728	NM_013217	v	
1626	1396	NM_013222	j	
1627	815	NM_013224	w	
1628	18305	NM_013226	V	Fatty acid metabolism, Propanoate
1629	21078	NM 016986	d	metabolism, Valine, leucine and isoleucine degradation, beta-Alanine metabolism
1630	24649	NM_016988	v	Riboflavin metabolism
1631	15239	NM_016989	q,w	
1632	45	NM_016996	General	
1633	20714	NM_016999	t	Fatty acid metabolism, Tryptophan metabolism Fatty acid metabolism, Tryptophan
1633	20713	NM_016999	t	metabolism
1633	20711	NM_016999	q,t	Fatty acid metabolism,Tryptophan metabolism
1633	20715	NM_016999	q,t	Fatty acid metabolism,Tryptophan metabolism
1634	1698	NM_017000	e,n,p,General	Sterol biosynthesis
				Glutathione metabolism,Pentose phosphate
1635	1399	NM_017006	h,n,General	cycle
1637 1638	18989 21013	NM_017013 NM_017014	n e.f	Glutathione metabolism Glutathione metabolism
1638	21015	NM 017014	e,General	Glutathione metabolism
1639	11836	NM 017023	b	
1639	5475	NM 017023	b	
1639	25546	NM_017023	b,bb	
1640	47907	NINA 047025	i,General	Cysteine metabolism,Glycolysis / Gluconeogenesis,Propanoate metabolism,Pyruvate metabolism
1640 1641	17807 24597	NM_017025 NM_017040	u	metabolism, Fyruvate metabolism
1642	24696	NM 017048	f,j,z	
1643	24695	NM_017049	u	
1644	20876	NM_017050	j,n,z	
1645	910	NM_017059	f,l,m	
1645	912	NM_017059	1	
1646	1946	NM_017061	h t General	
1646 1646	1942 1943	NM_017061 NM_017061	t,General t	
1647	6062	NM 017066	d	1
1648	6654	NM_017068	w	
1649	11153	NM_017073	S	Glutamate metabolism, Nitrogen metabolism
1650	923	NM_017076	General	
1651	1523	NM_017079	s	
1652	23660	NM_017080	s	Androgen and estrogen metabolism,C21- Steroid hormone metabolism
1653	275	NM_017081	b,d,General	Androgen and estrogen metabolism,C21- Steroid hormone metabolism
1654	16211	NM_017082	j,s,z	
1655	1552	NM_017084	j	Glycine, serine and threonine metabolism
1655	1550	NM_017084	у	Glycine, serine and threonine metabolism
1656 1657	22552 8888	NM_017087 NM_017090	a,k,x m	Purine metabolism
1658	10887	NM_017090 NM_017094	a,General	r unie nietabonsiii
1000	110001	114101_017034	ju, contrai	.L

TABLE 2: P	ATTHW/ANYS		ners alle sale	Aliy, Dodka: No. 44921-6089W
	1			Doc. No. 1798397.
ID No.	ldendder	ConBank Acel Ref. Seq. ID No.	Model Gode	Pathways
1659	4393	NM_017101	a,y	
1660	24770	NM_017111	d	
1661 1661	20745 20746	NM_017113 NM_017113	e a	
1662	1375	NM 017122	w	
1663	12903	NM_017124	k	
1664	24885	NM_017138	r	
1664	24886	NM_017138	d,q	
1665 1666	15363 13392	NM_017147 NM_017148	n,u u,General	
1667	5351	NM 017150	q	
1668	16954	NM_017151	a,n	
1669	21643	NM_017152	g	
1670	1694	NM_017153	a,q	
1671 1671	17104 17106	NM_017160 NM_017160	bb,General u	
1671	17106	NM 017160	d,e	
1672	17686	NM 017165	n,q	Glutathione metabolism
1673	20702	NM_017166	c	
1674	3513	NM_017177	r	Glycerolipid metabolism
1675	19031	NM_017180	v,General	
1676 1676	15437 15433	NM_017187 NM_017187	x,z v	
1676	15434	NM 017187	x,z	
1677	24437	NM 017190	p	
1678	1542	NM_017193	j,l,m,z	
1679	14695	NM_017202	q,s	Oxidative phosphorylation
1679	14694	NM_017202	s,z	Oxidative phosphorylation
1680	1428	NM_017213	m	
1681 1682	1622 13642	NM_017216 NM_017220	g,j,s,z v	
1682	19976	NM_017220	w	
1683	1510	NM_017224	General	
1684	1811	NM_017228	j,l,m,z	
1686	17563	NM_017245	a,c,e,q	
1687	17502	NM_017248	r	<u> </u>
1687 1688	17501 19	NM_017248 NM_017258	x v.General	
1689	15300	NM 017259	i,v,cc,General	
	1.5555		I,m,v,aa,cc,	
1689	15301	NM_017259	General	
1689	15299	NM_017259	I,y,cc,General	
1690	15224	NM_017264	d	Destace
1691 1692	3987 1447	NM_017280 NM_017281	bb	Proteasome Proteasome
1693	15535	NM_017283	s,bb	Proteasome
1694	12349	NM_017290	General	Oxidative phosphorylation
1695	15819	NM_017298	р	
1696	23825	NM_017299	V	
1696	23826	NM_017299	v	Older of the Charles
1697	14003	NM_017305	j,l,m,y,z	Glutamate metabolism, Glutathione metabolism
1698	26109	NM_017306	q,s	
1698	18687	NM_017306	q,t	Fatty acid metabolism
1699	18142	NM_017314	g,s,aa	
1700 1701	1894 20809	NM_017320 NM_017326	t u	
1701	355	NM_017334	cc	
1703	16148	NM_017340	q,s	Fatty acid metabolism
1703	16150	NM_017340	а	Fatty acid metabolism
1704	20849	NM_017343	r,u,General	
1704	20848	NM_017343	b,General	
1705	606	NM_017350	b	

VABUE 2: F	ATHWAYS			Atty, Dooret No, 44921-5089VVC Doe, No, 1798397.1
Sequence ID No.	ldenWfer	Cenbenk Acel Ref. Seq. ID No.	Modal Coda	Pathweys 🔾
1706	1581	NM_017365	General	
1707	455	NM 019131	x	
1707	456	NM 019131	y,z	
1708	4532	NM 019134	b	
1709	1608	NM 019166	j,y,z	
1710	7489	NM 019169	c,General	
1711	17066	NM_019170	р	Prostaglandin and leukotriene metabolism
1712	23924	NM_019174	bb	Nitrogen metabolism
1713	24019	NM_019186	t	
1714	22063	NM_019195	d	
1715	2079	NM_019220	j,k,z	
1716	16284	NM_019229	l,m	
1717	985	NM_019233	b,cc	
1718	15503	NM_019237	k,x	
1718	15504	NM_019237	k,x	
1719	17908	NM_019242	I,v,cc,General	
1720	11218	NM_019247	C	
1721	15259	NM_019259	d,f	
1722	21443	NM_019262	aa,General	
1722	21444	NM_019262	t,General	
1723	117	NM_019266	o,bb	
1724	1145	NM_019280	w	D'a contra de Caracia de Caracia
				Bile acid biosynthesis,Fatty acid metabolism,Glycerolipid metabolism,Glycolysis /
1725	22220	NM_019286	C	Gluconeogenesis, Tyrosine metabolism
1726	10015	NM_019289	I,m,t,x,General	
1726	10016	NM_019289	bb,General	
1727	21651	NM_019296	c,f,x	
1728	20751	NM_019301	S	
1729	645	NM_019345	bb	
1730	1301	NM_019349	C	
1731	3776 4592	NM_019354	a,u General	<u> </u>
1732 1733	1324	NM_019356 NM_019371	w	
1734	19577	NM 019377	e	
1735	24626	NM_019381	s	***************************************
1736	744	NM_019622	p	
1730	144	14141_019022	ΙΡ	Fatty acid metabolism, Tryptophan
1737	20716	NM_019623	c	metabolism
1738	20709	NM 019904	x	metabolism
1739	574	NM 019905	u,General	Glyoxylate and dicarboxylate metabolism
1740	9096	NM_019908	i	Cryoxylate and disarboxylate metabolism
1741	20457	NM 020073	i,General	
1741	20458	NM 020073	General	
1741	20460	NM 020073	General	
1742	18713	NM_020075	r	
1742	18715	NM 020075	r	
1743	20493	NM 020076	р	Tryptophan metabolism
1744	16375	NM_020976	g	
1745	20816	NM_021261	k,General	
1746	15335	NM_021264	а	
1747	18729	NM_021578	k,z	
1748	19060	NM_021587	cc	
1749	17324	NM_021593	o,General	
1750	19679	NM_021653	General	
1750	19678	NM_021653	a,v,General	
1751	19665	NM_021688	u,General	
1752	19667	NM_021690	m	
1754	22916	NM_021740	а	
1755	19710	NM_021744	t	
1755	19711	NM_021744	t	
1756	19712	NM_021745	Г	

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UANSILIE ZS IP			111111111111111111111111111111111111111	ANTY, Docket No. 44921-50397 Doc. No. 1793397
Sequence ID No	ldeniller	Cenbank Acel Ref. Seq. ID No.	Model Gode	Pathways 43
1757	1962	NM_021750	j,k,y,z	
1757	19824	NM_021750	a,bb	Taurine and hypotaurine metabolism
1758	25198	NM_021754	h .	
1758	20035	NM_021754	b,n,s,v,General	
1759	20090 17885	NM_021757 NM_021765	m	
1760 1762	20161	NM 021836	cc,General	
1764	1203	NM 021997	k,z	
1765	23151	NM 022005	b	
1767	17101	NM_022179	bb	Aminosugars metabolism,Fructose and mannose metabolism,Galactose metabolism,Glycolysis / Gluconeogenesis,Starch and sucrose metabolism Aminosugars metabolism,Fructose and mannose metabolism,Galactose
1767	17100	NM_022179	bb	metabolism,Glycolysis / Gluconeogenesis,Starch and sucrose metabolism
1768	20257	NM_022180	w,General	
1768	25699	NM_022180	li	
1768	10860	NM_022180	p	
1769	23780	NM_022183	k,x	
1770 1771	20312 6585	NM_022224 NM_022266	0	
1772	17161	NM 022298	d,p,cc i,v,cc,General	
1772	17162	NM 022298	u	
1772	17160	NM 022298	u	
1772	17158	NM_022298	q	
1773	11454	NM_022381	i,aa,General	
1773	11455	NM_022381	I,General	
1774	13480	NM_022390	S	Folate biosynthesis
1775	15184	NM_022391	z	
1776	22413	NM_022392	h	
1776 1777	22414 22499	NM_022392 NM_022393	n t	
1779	24537	NM 022399	e	, — 1, — 1, — 1, — 1, — 1, — 1, — 1, —
1779	24539	NM 022399	v	
1780	1141	NM 022401	o,General	
1781	1069	NM 022402	g	
1782	8211	NM_022500	j,n,s	
1782	8212	NM_022500	n,s	
1783	6815	NM_022503	s	Oxidative phosphorylation
1784	4259	NM_022504	q,w	
1785	2236	NM_022509 NM_022512	J V Z	Butanoate metabolism,Fatty acid metabolism,Valine, leucine and isoleucin degradation
1787	3026	NM 022514	y,z a	- Cognition
1787	3027	NM_022514	a,q,r,aa	
1788	2696	NM_022515	a,d	
1788	2697	NM_022515	n,w,aa	
1789	3900	NM_022516	h	
1790	4151	NM_022518	0	Arginine and proline metabolism, Urea cy
1791	4242	NM_022521	c	and metabolism of amino groups
1792	4412	NM_022523	0	
1793 1794	8097	NM_022533	General	
1795	8597	NM_022536 NM_022538	a c,r,u	-
1795	8598	NM 022538	u	
1796	9296	NM_022541	0	
1797	21063	NM_022585	h	

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TABLE 2:	PATHWAYS			(Alix Docket No. 44921 5089)
				Doc. No. 1798397
Bequence D No.	relilinebl.	GenBenk Acc Ref. Seq. ID No. :	Model Code	Pathways
799	20781	NM_022591	z	
800 801	20803	NM_022592 NM_022594	n	Carbon fixation, Pentose phosphate cycle
802	20925 20944	NM_022597	q aa	
803	21024	NM 022599	o,General	
804	2250	NM_022643	General	
805	17567	NM_022672	a,y	
806 807	17661 24563	NM_022674 NM_022676	bb b	
807	24564	NM 022676	b,x	
808	20506	NM_022686	l l	
809	20508	NM_022688	g	
810	17586	NM_022694	k	
811 811	17730 17729	NM_022697 NM_022697	а	
812	154	NM 022849	lt l	
813	127	NM_022855	h	
814	152	NM_022858	j	
816	18101	NM_022948	z	
816 817	18103 21491	NM_022948 NM_022951	w	
818	15742	NM 022958	lv	
819	9286	NM 023027	t,w	
820	23215	NM_023102	z	
821	21238_	NM_024125	cc,General	il6,interact6-1
821	21239	NM_024125	cc,General i,n,General	il6,interact6-1
822 822	353 354	NM_024127 NM_024127	i,n,General	
822	352	NM 024127	h,General	
823	17227	NM_024131	x	
824	1598	NM_024134	l	
825	1162	NM_024153	d	Porphyrin and chlorophyll metabolism Oxidative phosphorylation, Type III proteir
1826 1827	7863 22079	NM_024156 NM_024157	x	secretion system
828	16476	NM 024162	General	
829	17765	NM_024351	b,s,v	
1830	8879	NM_024360	h	
831	20772	NM_024363	x	
1832	2812	NM 024386	c	Butanoate metabolism, Synthesis and degradation of ketone bodies, Valine, leucine and isoleucine degradation
833	335	NM_024387	j,y	Porphyrin and chlorophyll metabolism
834	21	NM_024388	cc	
834	22	NM_024388	f	Androgen and estrogen metabolism
836 837	9929 3582	NM_024392 NM_024396	aa	Androgen and estrogen metabolism
838	19993	NM_024398	e,p,s,aa	
839	10789	NM_024399	0	
840	22626	NM_024400	cc,General	
841	13633	NM_024403	g,General	
841 842	13634 23387	NM_024403 NM_024404	g,General b,General	
843	21038	NM 024484	h	Glycine, serine and threonine metabolism
844	1853	NM_030826	s	Glutathione metabolism
845	15111	NM_030827	e,General	
845	15112	NM_030827	y,z	
845	15110	NM_030827	General	
846 847	808 4057	NM_030837 NM_030844	k,m k	-
848	1221	NM_030845	t	
849	21509	NM_030847	×	
1850	1928	NM_030872	v	

vabue2: P				Atty, Docket No. 44924-5089/ Doc. No. 1798397	
Sequence ID No.	ldeniiier '	Cenbank Acel Ref. Seq. ID No.'	Model Gode	Pathways . I	
1851	17342	NM_030873	u		
1852	24648	NM_030985	u		
1852	25453	NM_030985	General		
1853	21802	NM_030987	h		
1854	23109	NM_031000	f,s,z	Glycerolipid metabolism, Glycolysis / Gluconeogenesis, Pentose and glucurona interconversions	
1855	134	NM_031003	a,u		
1856	25461	NM_031009	0		
1857	1845	NM_031010	t	Prostaglandin and leukotriene metabolism	
1857	25517	NM_031010	c,t	Prostaglandin and leukotriene metabolisr	
1858	16562	NM_031020	f		
1859	1480	NM_031021	f		
1860	1719	NM_031024	n		
1861	1350	NM_031030_	h		
4860	40775	NINA 024024	Conoral	Arginine and proline metabolism, Glycine, serine and threonine metabolism, Urea cy and metabolism of amino groups	
1862	16775	NM_031031	General	and metabolism of amino groups	
1863 1864	691 15886	NM_031034 NM_031035	z		
1866	3608	NM 031044	k,General	Histidine metabolism	
1866	3610	NM 031044	d,General	Histidine metabolism	
1867	15137	NM 031051	s	Thistidine metabolism	
1868	514	NM 031056	General		
1008	514	NW_031030	General	Inositol metabolism,Propanoate metabolism,Valine, leucine and isoleucin	
1869	17269	NM_031057	General	degradation	
1870	11849	NM_031065	а		
1871	1855	NM_031074	h		
1872	4683	NM_031083	d		
1873	15202	NM_031093	a		
1873	15201	NM_031093	a,n		
1874	12639	NM_031099	aa		
1875	20812	NM_031100	a		
1876	16938	NM_031103	w		
1877	19268	NM_031104	q		
1878	16929	NM_031108	9		
1879	10878	NM_031110	q,bb		
1880	19162	NM_031111	aa		
1880	19161	NM_031111	a,bb		
1881	24615	NM_031112	а,у		
1882	20839	NM_031113	a,q		
1883	19040	NM_031114	I,m,General		
1884	16349	NM_031115	u General		
1885 1886	14970 1814	NM_031127 NM_031134			
1887	13359	NM_031135	n,q General	-	
1888	15052	NM_031136	a		
1888	19359	NM_031136	a		
1889	15185	NM 031140	General		
1890	21625	NM_031144	a,e		
1891	238	NM 031152	bb		
1891	240	NM_031152	bb		
1892	15277	NM_031237	9		
1893	18083	NM 031315	q		
1893	1858	NM 031315	q		
1894	15663	NM_031318	General		
1895	1422	NM_031324	bb,General		
				Nucleotide sugars metabolism,Pentose a glucuronate interconversions,Starch and	
1896	18597 11259	NM_031325	g,bb i,cc,General	sucrose metabolism	
1897		NM 031327			

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TABUE 28 P	ZYAWHTA			Atiy. Docket No. 44921-5089W0 Doc. No. 1793397.1
D No.	refilirebl	CenBank Acel Ref. Seq. ID No.	Model Gode	Pathways
1899	18375	NM_031331	l,m	
1900	3519	NM_031334	cc	
1901	20698	NM_031357 NM_031509	b	Glutathione metabolism
1903	634 25525	NM_031509	n n	Glutathione metabolism
1903	25069	NM 031509	b,n,w	Clatatione metabolism
1903	635	NM 031509	z	Glutathione metabolism
1904	848	NM_031517	t	
1905	1872	NM_031523	а	
1905	16245	NM_031523	a,d,u	
1905	16244	NM_031523	а	
1906	9370	NM_031527	W	
1907	20448	NM_031530	General General	
1907	20449	NM_031530	General	Androgen and estrogen
1908	14633	NM_031533	u	metabolism,Pentose and glucuronate interconversions,Porphyrin and chlorophyll metabolism,Starch and sucrose metabolism
1909	16048	NM_031541	f	'
1910	4011	NM_031543	c,q	Fatty acid metabolism, Tryptophan metabolism Fatty acid metabolism, Tryptophan
1910	4010	NM_031543	c,q	metabolism Fatty acid metabolism,Tryptophan
1910	4012	NM 031543	lq	metabolism
1911	28	NM_031546	General	
1912	24640	NM_031548	h,cc	
1913	17149	NM_031549	x	
1913	17151	NM_031549	x	
1914	13105	NM_031552	w	Fatty acid metabolism,Glycerolipid
1915	15411	NM 031559	d,r	metabolism
1916	16164	NM 031563	a,y	THE (ADDITION
1917	9621	NM 031570	bb	
1917	9620	NM_031570	w,bb	
1918	546	NM_031573	f	
1919	1921	NM_031576	f	
1919	1920	NM_031576	r	
1920	24219	NM_031579	i,General	
1921	770	NM_031584	k,x	
1922	18008 18005	NM_031588 NM_031588	cc h	
1922 1922	18005	NM_031588 NM_031588	cc,General	
1922	1584	NM_031595	k	
1924	24235	NM_031614	v	Pyrimidine metabolism
1924	24234	NM_031614	General	Pyrimidine metabolism
1925	1639	NM_031627	j,l,v	
1926	1727	NM_031642	m,General	
1927	20766	NM_031643	у	
1929	1993	NM_031655	k,I,m,General	
1930	2057	NM_031660	e l: Connect	
1931	15039	NM_031672	k,General	Butanoate metabolism,Fatty acid biosynthesis (path 2),Fatty acid metabolism,Lysine degradation,Tryptophan metabolism,Valine, leucine and isoleucine
1932	15175	NM_031682	bb	degradation
1933	1004	NM_031685	V	
1934	19727	NM_031687	a,q,s	
1935	20404	NM_031700	j,r,y	
1935	20405	NM_031700	lo,r	<u> </u>

TABLE 2: P				ATTY, Docker No. 44994-5039Wo Doc, No. 1793397.1
Sequence. ID No.	ldentiiter	Cenbank Acel Ref. Seq. (D No. W	Model Gode	
4000	044	NINA 024705	Consent	Pantothenate and CoA biosynthesis,Pyrimidine metabolism,beta- Alanine metabolism
1936	811	NM_031705	General	Pantothenate and CoA biosynthesis,Pyrimidine metabolism,beta-
1936 1937	812 16204	NM_031705 NM_031706	o,v,bb,General	Alanine metabolism
1937	16205	NM 031706	a,y	
1938	24081	NM 031708	m	
1939	16918	NM_031709	a,q	
1940	1081	NM_031712	General	
1941	1340	NM_031715	b,n,u,cc, General	Fructose and mannose metabolism,Galactose metabolism,Glycolysis / Gluconeogenesis,Pentose phosphate cycle
				Arginine and proline metabolism,Ascorbate and aldarate metabolism,Bile acid biosynthesis,Butanoate metabolism,Fatty acid metabolism,Glycerolipid metabolism,Histidine metabolism,Lysine degradation,Propanoate metabolism,Pyruvate metabolism,Tryptophan metabolism,Valine, leucine and isoleucine degradation.beta-
1942	23884	NM_031731	j,s	Alanine metabolism
1943	10241	NM 031740	d	
1944	1214	NM 031741	r	
1944	1215	NM 031741	r	
1945	20724	NM 031753	h	
1946	20753	NM 031763	h	
1946	20752	NM 031763	ly	
1947	14953	NM_031774	р	
1948	14184	NM 031776	t,General	Purine metabolism
1948	14185	NM 031776	d,o,t,General	Purine metabolism
1949	1169	NM_031789	С	
1950	16155	NM_031810	d,z	
1950	16156	NM_031810	d	
1951	17194	NM_031814	z	
1952	17535	NM_031816	bb	
1953	2655	NM_031821	i,I,m,aa	
1954	10167	NM 031830	li	
1955	22321	NM 031832	o,t,u,General	
1956	4748	NM 031834	e,t	
1956	4749	NM_031834	e,t	
	1	<u> </u>		Alanine and aspartate metabolism, Glycine,
1957	7914	NM 031835	е	serine and threonine metabolism
1958	8385	NM 031836	h	
1958	8384	NM_031836	h	
1959	10268	NM_031838	а	
1959	10269	NM_031838	aa	
1959	10267	NM 031838	n,aa	
1960	15077	NM_031841	b	
1961	16726	NM_031855	x	Fructose and mannose metabolism
1962	25802	NM_031969	a	
1962	19191	NM 031969	c	
1962	19195	NM 031969	r	
1962	19190	NM 031969	p	
1963	17734	NM 031970	v,General	
1964	1475	NM_031970	v,Gerierar	
1965	15470	NM_031978	f	
1966	18502	NM_031984	c	
1967	19768	NM 031986	v,aa,General	
1968	723			
1300	1123	NM_032084	lu	<u> </u>

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VABUE 2: PA	ATHWAYS			/Yiy. Dooket No. 44921-5089 Doc. No. 179339
Sequence D No.	ldeniiier	GenBenk Acel Ref. Seq. ID No.	Model Gode	Pathways N
1969	17935	NM_032615	а	
1970	16831	NM_033095	n	
1971 1971	25468 25469	NM_033234 NM_033234	c,z c	
1971	17832	NM 033234	c,p	
1971	17829	NM 033234	c,z	
1972	4723	NM_033235	z	
1973	1409	NM_033349	p,General	Pyruvate metabolism
1974	19998	NM_033352	General	
1975	1410	NM_052798	d	
1976	15028	NM_052809	f	Cysteine metabolism, Taurine and hypotaurine metabolism
1977	5176	NM_053297	u	
1978	7660	NM_053299	1	
1979 1981	5117 17473	NM_053310 NM_053319	p a,v	
1981 1982	25480	NM_053319 NM_053329	g g	
1982	21977	NM 053329	v	
1983	14926	NM 053330	f	
1983	14929	NM_053330	e,General	
1984	16407	NM_053332	c,e	
1985	15790	NM_053341	j,x	
1986	6154	NM_053356	р	
1987	9215	NM_053374	li .	
1988	6416	NM_053380	General	
1989 1990	19113 2242	NM_053395 NM_053433	n,General	
1990	5561	NM 053438	v	
1992	14670	NM 053439	n,General	
1993	17102	NM 053440	w	
1994	24762	NM_053442	General	
1995	8085	NM_053453	General	
1996	4622	NM_053463	d	
1997	21866	NM_053472	р	
1998	9573	NM_053475 NM_053480	h	
1999 2000	16137 15556	NM 053480	k v	
2000	16394	NM_053485	General	
2002	4290	NM 053487	j,y	
2004	18826	NM 053523	d	
2005	7764	NM_053525	aa	
2006	14199	NM_053538	С	
2007	1058	NM_053539	c,d	
2008	4327	NM_053563	General	
2009	1342	NM_053573	h	Methane metabolism,Phenylalanine
2010	19254	NM_053576	h,s	metabolism Methane metabolism,Phenylalanine metabolism
2010 2011	19253 3049	NM_053576 NM_053582	h p,cc,General	metabolism
2011	3050	NM 053582	o,General	
2012	21423	NM 053586	s,y	Oxidative phosphorylation
2013	21445	NM_053587	t,v	
2014	20871	NM_053591	j,l	
2014	20870	NM_053591	1	
2015	21044	NM_053594	d	
2016	21709	NM_053596	k	
2016 2017	21708 1597	NM_053596 NM_053611	z I+	
2017	5565	NM 053618	General	
2019	13004	NM_053623	1	Fatty acid metabolism

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				Attiy, Docket No. 44921-5039W0 Doc. No. 1793397.1
Sequence ID No.	ldentifer	GenBank Acal Ref. Seq. (D No.)	Model Gode	Pathways
				Arginine and proline metabolism, D-Arginine and D-ornithine metabolism, Glycine, serine
2020	1127	NM_053626	g	and threonine metabolism
2021	18644	NM_053648	n	
2022	21637	NM_053653	р	
2023	3454	NM_053662	cc	
2024	16121	NM_053698	h,j,z	
2024 2025	16122	NM_053698	h,j,z General	
2025	25379 13622	NM_053713 NM_053713	General	
2026	15376	NM 053747	h	
2027	1218	NM 053748	b	
2028	1137	NM 053763	v	
2029	15996	NM 053769	cc	
2030	8652	NM 053774	g	
2031	14664	NM 053806	General	
2032	4361	NM 053812	k	
2034	15002	NM 053819	b,x,bb,General	
			b,l,x,bb,	
2034	15003	NM 053819	General	
2035	16173	NM_053822	t	
2036	17154	NM 053835	j,z	
2037	20868	NM_053843	t	
2037	20869	NM_053843	t	
2040	714	NM_053863	у	
2041	19781	NM_053883	b	
2041	19780	NM_053883	b	
2042	1454	NM_053887	General	
2043	1660	NM_053891	g	
2044	712	NM_053896	k	
2045	753	NM_053897	k	
2046	794	NM_053902	General	Tryptophan metabolism
2047	17937	NM_053911	T	
2048	8188	NM_053927	General	
2050	1628	NM_053936	h	
2051 2052	13954 408	NM_053955 NM_053961	General General	
2052	19991	NM 053961	a	
2052	16190	NM 053961	q	
2052	21355	NM 053961	j,l,y,z	
2055	15136	NM 053971	aa	
2055	15135	NM 053971	d	***
2056	1764	NM 053974	h	
2057	1292	NM 053980		
2058	15468	NM 053982	q	
2059	15642	NM_053985	General	
2060	21066	NM_054001	t	
2061	17326	NM_054008	0	
2061	17327	NM_054008	cc	
2061	17329	NM_054008	g,o,cc	
2062	25253	NM_057099	j,l,m,p,z	
2062	22849	NM_057099	j,l	
2063	19657	NM_057103	b,cc	
2064	5492	NM_057105	w	Androgen and estrogen metabolism,Pentose and glucuronate interconversions,Porphyrin and chlorophyll metabolism,Starch and sucrose metabolism
2064	15126	NM_057105	r	Androgen and estrogen metabolism,Pentose and glucuronate interconversions,Porphyrin and chlorophyll metabolism,Starch and sucrose metabolism

TABUE 2: F	PAYTHWAYS			Aliy, podket No. 44921-5089W0 Doc. No. 1793397.1
Sequence [D] No.	ldeniller	Cenbank Ace Ref. Seg. ID No.	Model Gode	Pathways
				Androgen and estrogen
				metabolism,Pentose and glucuronate
2064	15125	NIM 057105		interconversions, Porphyrin and chlorophyll metabolism, Starch and sucrose metabolism
2066	15125 15391	NM_057105 NM_057114	n	metabolism, Starch and sucrose metabolism
2067	727	NM 057123	m	
2068	915	NM_057124	S	
2069	15151	NM_057131	k	
2070	1892	NM_057144	b	
2071	12333	NM_057155	t Conord	
2071 2071	12331 12332	NM_057155 NM_057155	v,General f,General	<u> </u>
2072	17477	NM_057194	a,General	
2073	15408	NM 057197	p,t	
2073	15409	NM_057197	t	
2074	7866	NM_057198	h	Glutamate metabolism,Purine metabolism
2075	14125	NM_057208	h,j,y,z	
2076	1743	NM_057210	k,s	
2077	10498	NM_078617 NM_080399	a	
2078 2079	8820 15701	NM 080581	n j,m,y,z	
2079	20105	NM 080581	aa	
2080	16109	NM 080585	c	
2081	1757	NM 080766	d	
2082	7108	NM_080778	У	
2083	132	NM_080782	k	
2083	133	NM_080782	1	
2084	20122	NM_080887	General	
2085	6143	NM_080892	le lh	
2086 2087	9952 17546	NM_080902 NM_130401	b	
2088	21695	NM 130411	c,x	
2089	21391	NM_130416	x,General	
2090	20694	NM_130430	General	
2090	19818	NM_130430	СС	
2090	18810	NM_130430	e,s	
0004	10202	NINA 420422		Bile acid biosynthesis,Fatty acid biosynthesis (path 2),Fatty acid metabolism,Phenylalanine metabolism,Valine, leucine and isoleucine degradation
2091	18293 25064	NM_130433 S45392	q a,n	degradation
2093	3244	S63519	u	
2094	25501	S63521	q	
2095	16248	S68135	h	
2096	18647	S69316	q	
2097	24351	S74257	V	
2098	25066	\$75280	j,l,m,x,y,	
2099	1460	S76054	General	
2100	25539	S76742	V	
2101	16400	S76779	C	
2102 2103	24469 25545	S77858 S77900	h k,s	
2103	21583	S77900 S77900	k	
2104	10260	S81497	s	
2105	3609	S82579	k	Histidine metabolism
2106	111	U02506	u	
2107	14959	U03390	a,q,General	
2109	2010	U05675	b,x,bb	
2110	15462	U06230	d Conoral	
2112	1583	U07201	s,General	
2113	627	U09229	<u> h</u>	

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TABLE 2: F			4	ATTY, 1000 Fet 11/0, 44/92/1-5089WG Doc. No. 1798397.1
D No.	ldemilier	Consenk Acel Ref. Seq. ID No.	 	Pathways 4
2114	809	U17035	General	
2115	16675	U17565	k,x,bb	
2116	25587	U20110	Г	
2117	90	U20796	r b oo	
2118 2119	25589 22196	U21718 U21719	h,aa h	
2120	17118	U25746	s	
2121	1537	U27518	g,h,n	
2122	1558	U28504	bb	
2123	16193	U30831	n	
2124	17480	U31598	z	
2125	18302	U33500	General	
2126	25599	U34897	У	
2127	1394	U37099	h	
2128	244	U38376 U41164	ln lh	
2129 2130	1623 15851	U41164 U42719	f,t,x,General	
2130 2131	17886	U47315	s,z	
2132	21654	U53184	i,t,General	
2133	1439	U57391	w	
2134	725	U62316	bb	
2137	2153	U75404	b,cc,General	
2139	4956	U76714	j,y	
2140	4477	U77829	l,m	
2141	21703	U82591	z	
2142	977	U89744	s	
2143 2144	23282 22005	U90725 U96490	lm	
2146	819	X02284	j,z	Carbon fixation, Fructose and mannose metabolism, Glycolysis / Gluconeogenesis, Inositol metabolism, Pentose phosphate cycle Carbon fixation, Fructose and mannose metabolism, Glycolysis / Gluconeogenesis, Inositol
2147	818	X02291	e,j,z	metabolism,Pentose phosphate cycle
2148	20818	X02904	n,q	Glutathione metabolism
2149	16401	X04979	c	
				Carbon fixation, Glycolysis /
2450	20513	X05684		Gluconeogenesis,Purine metabolism,Pyruvate metabolism
2150 2151	25084	X06769	o,r cc	metabolism, Fyruvate metabolism
2152	672	X13722	h	1
2153	25675	X14181	n	
2153	20810	X14181	n,q,w	
2154	18541	X14671	у	
2155	25679	X15013	q	
2155	19244	X15013	c,q,w	
2156	15626	X17665	а	
2157 2158	1893 25686	X51529 X51536	t bb	Glycerolipid metabolism,Phospholipid degradation,Prostaglandin and leukotriene metabolism
2158	10819	X51536	aa,bb	
2159	18250	X51706	a,q,w	
2160	20872	X51707	а	
2161	516	X52711	С	
2162	25689	X52815	9	
2163	20427	X53378	W Conord	
2164 2165	18606 1463	X53504 X54467	General d,u,General	
2166	24577	X55153 ·	a,v	
			1 -4 1 7	

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TABLE 23: P	AVIHWAYS	processing of the processing of the contract o		/Xiiy, Docket No. 44921-5039W0 Doc. No. 1793397.1
Sequence ID No.	refillmebl	GenBank Acc/ Ref. Seq. ID No.	Model Gode	
2168	15106	X57529	g,n,q	
2169	5667	X58200	q,bb	
2169	18611	X58200	a,v	
2170	17175	X58389	w	
2171	25702	X58465	w	
2171 2172	10109 25705	X58465 X59375	c,q c,i,aa,General	
2173	25709	X59737	u	
2174	18354	X59859	General	
2174	18355	X59859	t	
2175	21657	X61381	General	
2176	25718	X62145	bb,General	
2176	15875	X62145	a,q,v	
2177	13646	X62166	bb	
2178	25721	X62325	р	
2179	16012	X62875	m,s,z	
2180	25730	X63369	cc	
2181	25089	X63594	General	
2181	25090	X63594	cc,General	
2182	20844	X65228	n,w	
2183 2184	20879 25736	X65296 X68782	j,y c	
2185	16426	X70369	c	
2186	16300	X70706	u	
2187	24232	X75207	C	
2188	16272	X76456	n,p	
2189	25741	X76489	u	
2190	23302	X78949	h	
2191	25747	X81448	General	
2192	24115	X81449	u	
2193	25754	X89696	g	
2194	25097	X90642	y,z	
2195	12978	X96437	cc,General	
2197	4594	Y07704	С	
2198	25777	Y08355	g,p,General	
2199	15986	Y09945	bb,General	
2200	20890	Y13275	k	
2201	21914	Y13336 Z11995	d Coporal	
2202 2203	406 18352	Z12298	o,General t	
2204	17481	Z49761	k	
2205	8664	Z75029	r,v	
2206	2459	AA964755	cc	
2207	23830	AA956638	aa	
2208	6100	X73524	x	
2209	439	Z22607	w	
2210	8665	AI071965	v	
2211	155	U32681	t	
				Methane metabolism,Phenylalanine
2212	19252	AA892041	s	metabolism
2213	15582	AI232320	q	
2214	17541	M26125	n	
2215 2216	18609	M30689	i	
2216	6262 23859	AI177125 AI072161	g f	
2218	21011	H32189	e	Glutathione metabolism
2220	2572	Al177143	b	Gratatilone metabolism
2221	25419	M22922	a	

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TABLES		HOMOLOGUEAN	INOTATIONS		Auy, Docket No. 44921-5039WÖ
					Doc. No. 1793397.1
809. ID	lemilier	Consonk Ace <i>l</i> Ref. Seq. ID No.	Model Code	Kamalogovs Cana	Homologous Christer Namo
1	6949	AA012785	q		
2	25098	AA108277	h,v		EST, Moderately similar to A Chain A, Mdm2 Bound To The Transactivation Domain Of P53 (SUB 17-125 [H.sapiens], mouse double minute 2, human homolog of; p53-binding
3	17312	AA108308	r		protein NADH dehydrogenase (ubiquinone) 1
4	16882	AA684537	0		beta subcomplex, 5 (16kD, SGDH)
5 6	6049 4426	AA685178 AA685974	y I.m		EST, Weakly similar to T30827 nascent polypeptide-associated complex alpha chain, non-muscle splice form - mouse [M.musculus], FKSG17, Homo sapiens alpha-NAC gene for nascent polypeptide- associated complex component, KIAA0363 protein, expressed sequence AL022831, nascent- polypeptide-associated complex alpha polypeptide
0	4420	AA003914	1,111		EST, Weakly similar to T46390
7	21815	AA686423	g		hypothetical protein DKFZp434C1920.1 [H.sapiens], hepatocellular carcinoma-associated antigen 59
8	1600	AA686470	i	DNA-damage inducible transcript 3, DNA-damage-inducible transcript 3	DNA-damage inducible transcript 3, EST, Moderately similar to GA15_HUMAN GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE PROTEIN GADD153 [H.sapiens], myozenin
8 9	1599 21997	AA686470 AA799325	i u	DNA-damage inducible transcript 3, DNA-damage-inducible transcript 3	DNA-damage inducible transcript 3, EST, Moderately similar to GA15_HUMAN GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE PROTEIN GADD153 [H.sapiens], myozenin
10	18396	AA799330	v		
			,		ESTs, Highly similar to ERR3_HUMAN ESTROGEN-RELATED RECEPTOR GAMMA [H.sapiens], Untitled, estrogen related receptor, alpha, estrogen related receptor, beta,
11	6581	AA799412	f,I		estrogen-related receptor beta
					ESTs, Moderately similar to NPL4_HUMAN NUCLEOSOME ASSEMBLY PROTEIN 1-LIKE 4 [H.sapiens], ESTs, Weakly similar to NPL4_HUMAN NUCLEOSOME ASSEMBLY PROTEIN 1-LIKE 4 [H.sapiens], SET translocation, nucleosome assembly protein 1-like 1,
12	16538	AA799449	k		nucleosome assembly protein 1-like 4
13	23294	AA799472	u		CGI-116 protein
14	18290	AA799497	r	<u> </u>	

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TABLE 8	S CHUMIAN	HONOLOGAE V	SKOJVATOKI		Atty: Docket No. 4/1924-5039WO
		CenBenk Ace./ Ref. Seq. ID No.		Mame Mame	Homologous Gluster Kame
					DAZ associated protein 1, ESTs, Highly similar to ROA1 RAT HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEIN A1 [R.norvegicus], ESTs, Moderately similar to Up1, The Two Rna-Recognition Motif Domain Of Hnrnp A1 (SUB 3-184 [H.sapiens], ESTs, Weakly similar to ROA1 RAT HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEIN A1 [R.norvegicus], ESTs, Weakly similar to ROA2 MOUSE HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEINS A2/B1 [M.musculus], RIKEN cDNA 3010025E17 gene, Ras-GTPase activating protein SH3-domain-binding protein 2, Ras-GTPase-activating protein SH3-domain binding protein, cell death regulator aven, heterogeneous nuclear ribonucleoprotein A1, heterogeneous
15	18981	AA799523	е		nuclear ribonucleoprotein A2/B1 EST, Moderately similar to A38983 TCP1 ring complex protein TRiC5 [H.sapiens], T-complex 1, chaperonin
16 17	20843 16993	AA799545 AA799560	h b		containing TCP1, subunit 3 (gamma), expressed sequence Al528772, t- complex 1, t-complex protein 1
18	16576	AA799570	d		
19	18361 17712	AA799591 AA799598	i Z		EST, Moderately similar to I38369 beta-tubulin [H.sapiens], ESTs, Highly similar to I38370 beta-tubulin [H.sapiens], ESTs, Highly similar to T08726 tubulin beta chain [H.sapiens], ESTs, Highly similar to TBB1 RAT TUBULIN BETA CHAIN [R.norvegicus], ESTs, Moderately similar to I38370 beta-tubulin [H.sapiens], RIKEN cDNA 2410129E14 gene, RIKEN cDNA 4930447K03 gene, RIKEN cDNA 4930542G03 gene, Rat mRNA for beta-tubulin T beta15, beta tubulin 1, class VI, tubulin, beta 3, tubulin, beta, 2
22	18346 8768	AA799718 AA799726	f		
24	11687 18349	AA799732 AA799744	w		DiGeorge syndrome chromosome region 6, DiGeorge syndrome critical region gene 6, DiGeorge syndrome critical region gene 6 like
26	17494	AA799751	n		
27	18360	AA799771	General		
28	20998	AA799803	z		EST, Moderately similar to MAS2_HUMAN MANNAN-BINDING LECTIN SERINE PROTEASE 2 PRECURSOR [H.sapiens], Rattus norvegicus mRNA for serine protease, complete cds, complement C1r-like proteinase precursor,, complement component 1, r subcomponent, complement component 1, s subcomponent, protein C
		1. 5 (1 55000	ı -		Teacomponent, protein o

TABUE	E HUMAN	HOMOROGAE W	BNOTATON		Aily, Docket No. 44921-5939W0 Doc. No. 1793897.1
Seg. ID No.	ldentifier	Conbank Ace <i>l</i> Ref. Seq. (D No.	Model Code	Homologous Cone Namo	Honologous Cluster Name
30	21006	AA799861	С		interferon regulatory factor 7
30	21000	701133331			ESTs, Moderately similar to Up1, The Two Rna-Recognition Motif Domain Of Hnrnp A1 {SUB 3-184 [H.sapiens], ESTs, Weakly similar to ROA1 RAT HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEIN A1 [R.norvegicus], ESTs, Weakly similar to heterogeneous ribonuclear particle protein A1 [H.sapiens], Human DNA sequence from clone RP11-51N22 on chromosome 13 Contains ESTs, STSs and GSSs. Contains an HNRPA1 (heterogeneous nuclear
					ribonucleoprotein A1) pseudogene, RIKEN cDNA 4930547K05 gene, heterogeneous nuclear
31	15011	AA799893	General		ribonucleoprotein A1
32 33	20811 23202	AA799899 AA799971	a General		EST, Moderately similar to RL1X_HUMAN 60S RIBOSOMAL PROTEIN L18A [H.sapiens], EST, Weakly similar to RL1X_HUMAN 60S RIBOSOMAL PROTEIN L18A [H.sapiens], EST, Weakly similar to S47353 ribosomal protein L18a, cytosolic [H.sapiens], ESTs, Highly similar to RL1X_HUMAN 60S RIBOSOMAL PROTEIN L18A [H.sapiens], RIKEN cDNA 2510019J09 gene, ribosomal protein L18a
33	23202	AA799971	General	-	ESTs, Moderately similar to 1701409A
34 35 36	4832 21656 18433	AA800190 AA800202 AA800218	b d j,y,z		glycogen phosphorylase [H.sapiens], ESTs, Weakly similar to 1701409A glycogen phosphorylase [H.sapiens], phosphorylase, glycogen; brain
37	6386	AA800235	u		
38	18442	AA800258	h,k		
39	21092	AA800380	у		DNA segment, Chr 14, University of California at Los Angeles 2, Hydroxysteroid dehydrogenase, 11 beta type 1, expressed sequence C79874, hydroxysteroid (11-beta) dehydrogenase 1, hydroxysteroid 11-beta dehydrogenase 1, hydroxysteroid 17-beta dehydrogenase 11, retinal short-chain dehydrogenase/reductase retSDR2
40	17325	AA800587	General		ESTs, Highly similar to GSHG_MOUSE GLUTATHIONE PEROXIDASE-GASTROINTESTINAL (GSHPX-GI) [M.musculus], Glutathione peroxidase 1, glutathione peroxidase 2 (gastrointestinal)
	40000	A A DOOC 4 O			zinc finger protein 36, zinc finger protein homologous to Zfp-36 in
41	13930	AA800613	cc, General		mouse
42 42	21372 21373	AA800693 AA800693	v s		
174	18161	AA800693 AA800701	k	-	
143		1 1000.01	1		1
43 44	6595	AA800753	w		

	10 10 10 10 10 10 10 10 10 10 10 10 10 1	HOMOLOGUE AN	RIGITATIONS		Any, Dooket No. 4/92/1-5089/W Dog. No. 17/93397.
Scq. (id) (No. : : ්	ldenilliler	GenBank Ace <i>l</i> Ref. Seq. ID No.	Model Code	Memologous Gene Name	Honologous Clyster Name
46	23115	AA801165	o,y		EST, Weakly similar to H2AL_HUMAN HISTONE H2A.L (H2A/L) [H.sapiens], H2A histone family, member L, similar to H2A histone family, member A (H. sapiens)
47	12399	AA801307	General		
48	7543	AA801395	General		
49	24237	AA817726	t,General		
50	11215	AA817921	0		
51	5985	AA818005	g		
52	11338	AA818016			EST, Weakly similar to RB6K MOUSE RABKINESIN-6 [M.musculus], RAB6 interacting, kinesin-like (rabkinesin 6), RIKEN cDNA 3110001D19 gene, Rab6, kinesin-like
52	11338	AA818016	x		COP9 (constitutive
					photomorphogenic), subunit 6 (Arabidopsis), Homo sapiens cDNA FLJ14833 fis, clone OVARC1001171, moderately similar to Homo sapiens translation initiation factor 3 47 kDa subunit mRNA, IFP38, RIKEN cDNA 0610037M02 gene, eukaryotic translation initiation factor 3, subunit 5 (epsilon, 47kD), hypothetical protein MGC13045, proteasome (prosome, macropain) 26S subunit, non-ATPase,
53	2845	AA818026	k,General		7
54	16756	AA818089	i,k,General		glycyl-tRNA synthetase
55	17771	AA818224	e,g,p,General		EST, Weakly similar to S45140 tubulin beta chain [H.sapiens], RIKEN cDNA 2410129E14 gene, RIKEN cDNA 4930542G03 gene, tubulin, beta 2, tubulin, beta 5, tubulin, beta polypeptide, tubulin, beta, 2
56	6522	AA818261	g,m		
57	5924	AA818359	у		
58	7806	AA818421	b,aa		
59	8237	AA818512	v		
60	17434	AA818574	h		
61	8728	AA818615	General		
60	5054	AA949355		diphtheria toxin receptor (heparin-binding epidermal growth factor-like growth factor), heparin binding epidermal growth factor-	diphtheria toxin receptor (heparin- binding epidermal growth factor-like growth factor), expressed sequence AW047313, heparin binding epidermal
62	6054	AA818658	b,v,cc,General	like growth factor	growth factor-like growth factor
63	11590	AA818721	a Conoral		
64 65	4291 4330	AA818741 AA818747	q,General o,General		
	19723	AA818747 AA818761	v,General	 	
66 67	13684	AA818770	h,j,l,m		
68	6322	AA818801	k	 	
00	0322	LV010001	IV.		expressed sequence AV066530, guanylate cyclase activator 2B (uroguanylin), guanylate cyclase
69	7690	AA818875	General		activator 2b (retina)
70	4952	AA818907	q,General		
71	6094	AA818911	t		
72	10985	AA818998	o,General		
73	6120	AA819008			

74 2586 AA8 76 6438 AA8 77 24721 AA8 78 6250 AA8 80 6281 AA8 81 10141 AA8 82 6551 AA8 83 6723 AA8	A819306 A819376 A819517 A819526 A819558 A819653		Honologeus Gene	Bog. No. 1798397.1 Homologous Gluster Name CocoaCrisp, ESTs, Weakly similar to JC5308 testis-specific, vespid, and pathogenesis-related protein 1 precursor [H.sapiens], Homo sapiens, Similar to RIKEN cDNA 1700011E04 gene, clone MGC:26856 IMAGE:4822995, mRNA, complete cds, RIKEN cDNA 1200009H11 gene, RIKEN cDNA 1700011E04 gene, RIKEN cDNA 4921508O11 gene, acidic epididymal glycoprotein-like 1, glioma pathogenesis-related protein, specific granule protein (28 kDa), testis specific gene 1, testis specific protein 1, testis specific protein 1 (probe H4-1 p3-1) ESTs, Weakly similar to T17246 hypothetical protein DKFZp586M0617.1 [H.sapiens], KIAA0263 gene product, mammalian inositol hexakisphosphate kinase 2 HYA22 protein, conserved gene amplified in osteosarcoma, nuclear
74 2586 AA8 76 6438 AA8 77 24721 AA8 78 6250 AA8 80 6281 AA8 81 10141 AA8 82 6551 AA8 83 6723 AA8	A819081 A819269 A819306 A819376 A819517 A819526 A819558 A819653	c o d,w		CocoaCrisp, ESTs, Weakly similar to JC5308 testis-specific, vespid, and pathogenesis-related protein 1 precursor [H.sapiens], Homo sapiens, Similar to RIKEN cDNA 1700011E04 gene, clone MGC:26856 IMAGE:4822995, mRNA, complete cds, RIKEN cDNA 1200009H11 gene, RIKEN cDNA 1700011E04 gene, RIKEN cDNA 4921508O11 gene, acidic epididymal glycoprotein-like 1, glioma pathogenesis-related protein, specific granule protein (28 kDa), testis specific gene 1, testis specific protein 1, testis specific protein 1 (probe H4-1 p3-1) ESTs, Weakly similar to T17246 hypothetical protein DKFZp586M0617.1 [H.sapiens], KIAA0263 gene product, mammalian inositol hexakisphosphate kinase 2
76 6438 AA8 77 24721 AA8 78 6250 AA8 80 6281 AA8 81 10141 AA8 82 6551 AA8 83 6723 AA8	A819269 A819306 A819376 A819517 A819526 A819558 A819653	o d,w o,y j		ESTs, Weakly similar to T17246 hypothetical protein DKFZp586M0617.1 [H.sapiens], KIAA0263 gene product, mammalian inositol hexakisphosphate kinase 2 HYA22 protein, conserved gene
77 24721 AA8 78 6250 AA8 80 6281 AA8 81 10141 AA8 82 6551 AA8 83 6723 AA8	A819306 A819376 A819517 A819526 A819558 A819653	o,y j		hypothetical protein DKFZp586M0617.1 [H.sapiens], KIAA0263 gene product, mammalian inositol hexakisphosphate kinase 2 HYA22 protein, conserved gene
78 6250 AA8 80 6281 AA8 81 10141 AA8 82 6551 AA8 83 6723 AA8	A819376 A819517 A819526 A819558 A819653	o.y j		hypothetical protein DKFZp586M0617.1 [H.sapiens], KIAA0263 gene product, mammalian inositol hexakisphosphate kinase 2 HYA22 protein, conserved gene
80 6281 AA8 81 10141 AA8 82 6551 AA8 83 6723 AA8	A819517 A819526 A819558 A819653	j j		HYA22 protein, conserved gene
81 10141 AA8 82 6551 AA8 83 6723 AA8	\819526 \819558 \819653	j j		
82 6551 AA8 83 6723 AA8	\819558 \819653	j t		LIM interactor-interacting factor
83 6723 AA8	\819653	t		
84 14958 AAE		r		
	\819744	aa		
	N 819776	v		ESTs, Highly similar to HS9B RAT HEAT SHOCK PROTEIN HSP 90-BETA [R.norvegicus], expressed sequence AL024080, expressed sequence C81438, heat shock 90kD protein 1, beta, heat shock protein, 84 kDa 1
86 6204 AA8	\819889	aa		0.19=0.1
	A848315	General		GMPR2 for guanosine monophosphate reductase isolog, IMP (inosine monophosphate) dehydrogenase 2, RIKEN cDNA 2310004P21 gene, RIKEN cDNA 5730544D12 gene, expressed sequence AA959850, guanosine monophosphate reductase, inosine 5'-phosphate dehydrogenase 2
	\848389	bb		
89 21125 AA8	\848437	General		
90 23504 AA8	\848496	q		ESTs, Moderately similar to IF4B_HUMAN EUKARYOTIC TRANSLATION INITIATION FACTOR 4B [H.sapiens], eukaryotic translation initiation factor 4B
		g		ESTs, Highly similar to FMO2_HUMAN DIMETHYLANILINE MONOOXYGENASE [H.sapiens], Flavin-containing monooxygenase 1, flavin containing monooxygenase 1, flavin containing monooxygenase 2, flavin containing monooxygenase 3, hypothetical protein PRO1257
		С		
		0		
		Gonoral Conoral		
	\849497 \849518	General I		
		i,I,m,General		

TABLE 8	B: HUMAN	HOMO! OGNE W			Affy, Docket No. 44921-5039000 Doc. No. 1793397.
809. (D No.	ldentifier	ConBenk Acel Ref. Seq. ID No.	Model Gode 🚧	Homologous Cento Namo	Homologous Cluster Name
98	16678	AA849827	aa		
99	8515	AA849917	е		
100	18447	AA849939	General		
101	12130	AA850037	р		
102	23981	AA850040	x,aa		RIKEN cDNA 2810452G09 gene, adenylyl cyclase-associated CAP protein homolog 1 (S. cerevisiae, S. pombe), adenylyl cyclase-associated protein
103	13615	AA850364	τ		DVEZD 40 40 40 5 manhair
105	2637	AA850893	×		DKFZP434O125 protein
106	22093	AA850909	d		
	21766	AA850916	C		ESTs, Highly similar to LB4D_HUMAN NADP-DEPENDENT LEUKOTRIENE B4 12-HYDROXYDEHYDROGENASE [H.sapiens], ESTs, Weakly similar to FAS RAT FATTY ACID SYNTHASE [R.norvegicus], ESTs, Weakly similar to LB4D_HUMAN NADP-DEPENDENT LEUKOTRIENE B4 12-HYDROXYDEHYDROGENASE [H.sapiens], crystallin, zeta, fatty acid
108	2847	AA850919	w		synthase
109	12162	AA850975	h		
110	9514	AA850978	General		
111	3924	AA851017	e,q		
111	3925	AA851017	o,General		
112	4490	AA851184	a.k		cathepsin Z
113	19187	AA851230	General		
114	19189	AA851237	С		RIKEN cDNA 1110058H21 gene, ubiquitin specific protease 18
115	15386	AA851241	m		breast cancer metastasis-suppressor 1, hypothetical protein MGC11296
116	21462	AA851261	g,I,General		ART-4 protein
117	21471	AA851343	General		NADH dehydrogenase (ubiquinone) Fe S protein 8 (23kD) (NADH-coenzyme
118	16902	AA851379	p		Q reductase)
119	23376	AA851392	i,x		kinesin-like 4
119	23377	AA851392	x		kinesin-like 4
120	13349	AA851417	General		
121	21527	AA851733	r,u		
122	4048	AA851814	i,o,u,General		EST, Moderately similar to PM17 MOUSE MELANOCYTE PROTEIN PMEL 17 PRECURSOR [M.musculus] Homo sapiens, Similar to glycoprotein (transmembrane) nmb, clone MGC:1696 IMAGE:3345861, mRNA, complete cds, glycoprotein (transmembrane) nmb, silver
					signal sequence receptor, alpha
123	10561	AA851871	bb		(translocon-associated protein alpha) ESTs, Weakly similar to A60021 tropomyosin-related protein, neuronal- rat [R.norvegicus], RIKEN cDNA 0710005K15 gene, expressed sequence R75279, reticulon 1,
124	17411	AA858621	j,y		expressed sequence Al747533, mini chromosome maintenance deficient 7 (S. cerevisiae), minichromosome
125	1801	AA858636	k,s,x,bb		maintenance deficient (S. cerevisiae)
126	18350	AA858674	р		
127	19484	AA858693	е		
128	6360	AA858696	d		

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ĪABUES: β		HONOLOGAL	RIOTETON		Ally, Dookel No. 44921-5039WO Doc. No. 1793397.1
Seq. [D]	141	Consont Acc.		Hamalogous Gene	
No. I di	entitier	Ref. Seq. ID No.	Model Gode	Namo 🐫 💮 🖂 🖂	Homologous Cluster Name
129 17:	334	AA858704	p		exostoses (multiple) 1, exostoses (multiple)-like 1, expressed sequence AA409028
130 63		AA858758	9		
		AA858759	a		
132 63	84	AA858788	I,m,General		
					LanC (bacterial lantibiotic synthetase component C)-like, LanC (bacterial lantibiotic synthetase component C)-
		AA858830	р		like 1, RIKEN cDNA 1700003F10 gene
135 72		AA858892	f		
136 18	217	AA858930	t		anners simul ADNIA systhators
137 580	67	AA858953	v,General		asparaginyl-tRNA synthetase, hypothetical protein FLJ23441 Interleukin 1 receptor accessory
138 14	479	AA858969	•		protein, Mus musculus IL-1Rrp2 mRNA, complete cds, interleukin 1 receptor accessory protein-like 2, interleukin 1 receptor, type I, interleukin 18 receptor 1
139 64		AA859085	t		Interiedkii 18 receptor 1
		AA859114	o.General		
		AA859241 AA859271	General c		EST, Highly similar to OM25_RAT MITOCHONDRIAL OUTER MEMBRANE PROTEIN 25 (NPW16) [R.norvegicus], EST, Weakly similar to OM25_RAT MITOCHONDRIAL OUTER MEMBRANE PROTEIN 25 (NPW16) [R.norvegicus], Erbb2 interacting protein, discs, large homolog 4 (Drosophila), expressed sequence Al118201, hypothetical protein FLJ11271, synaptojanin 2 binding protein
143 21	791	AA859333	k		EST, Moderately similar to CYSR RAT CYSTEINE-RICH PROTEIN 1 [R.norvegicus], ESTs, Weakly similar to S12658 cysteine-rich protein [H.sapiens], cysteine and glycine-rich protein 1, cysteine and glycine-rich protein 3 (cardiac LIM protein), cysteine rich protein, cysteine-rich protein 2, cysteine-rich protein 3, thymus LIM protein
			cc,General		
		AA859520 AA859545	f r		
					EST, Highly similar to SPERM-COATING GLYCOPROTEIN PRECURSOR [R.norvegicus], ESTs, Weakly similar to JC4131 glioma pathogenesis-related protein [H.sapiens], Human DNA sequence from clone RP5-881L22 on chromosome 20 Contains ESTs, GSSs, STSs and CpG islands. Contains a gene for a novel protein similar to a trypsin inhibitor and four other genes for novel proteins, RIKEN cDNA 1810049K24 gene, RIKEN cDNA 9230112K08 gene, acidic epididymal glycoprotein 1, acidic epididymal glycoprotein 2, epididymal glycoprotein, glioma pathogenesis-
47 198	394		AA859581	AA859581 s	AA859581 s

		HOWOTOGAL WK	SUCITATON		Affy, Docket No. 44921-5039W0 Doc. No. 1793397.1
S@, ID Ko.		ConBonk Ace./ Ref. Seq. ID No.		Homologous Gene Name	Homologous Chata Name
148	14353	AA859585	h		DnaJ (Hsp40) homolog, subfamily B, member 1, DnaJ (Hsp40) homolog, subfamily B, member 1, DnaJ (Hsp40) homolog, subfamily B, member 6, DnaJ (Hsp40) homolog, subfamily B, member 8, ESTs, Weakly similar to DnaJ-like protein [M.musculus], ESTs, Weakly similar to HS44 MOUSE HEAT SHOCK 40 KDA PROTEIN 4 [R.norvegicus], Homo sapiens cDNA FLJ13992 fis, clone Y79AA1002139, weakly similar to DNAJ PROTEIN HOMOLOG 1, RIKEN cDNA 1700029A20 gene, RIKEN cDNA
149	16318	AA859648	h		2010306G19 gene
150	17316	AA859652	General		
151 152	19067 22406	AA859663 AA859680	n,q n		
152	20599	AA859690	x		
154	14261	AA859693	ů	ļ	
155	14138	AA859700	v		protoporphyrinogen oxidase
155	14139	AA859700	v		protoporphyrinogen oxidase
157	22374	AA859804	1		
158 159 160	22385 22773 22816	AA859805 AA859885 AA859898	b,k n k,x,z		ESTs, Weakly similar to PROTEIN- LYSINE 6-OXIDASE PRECURSOR [R.norvegicus], Lysyl oxidase, lysyl oxidase, lysyl oxidase-like, lysyl oxidase-like 1, lysyl oxidase-like 2
161	11891	AA859926	X		
162	23070	AA859942	k		EST, Weakly similar to JC1343 glycylpeptide N- tetradecancyltransferase [H.sapiens], N-myristcyltransferase 1
163	23121	AA859948	k		
165	18468	AA859954 AA859966	cc,General aa		Homo sapiens cDNA FLJ14666 fis, clone NT2RP2003000, weakly similar to TUMOR NECROSIS FACTOR, ALPHA-INDUCED PROTEIN 1, Homo sapiens polymerase delta-interacting protein 1 mRNA, complete cds, MSTP028 protein, tumor necrosis factor, alpha-induced protein 1 (endothelial) ESTs, Moderately similar to A Chain A, Inositol Monophosphatase [H.sapiens], Inositol (myo)-1(or 4)-monophosphatase 1, RIKEN cDNA 2900059K10 gene, bisphosphate 3'-nucleotidase 1, inositol (myo)-1(or 4)-monophosphatase 1, inositol (myo)-
166	23336	AA859981	q		1(or 4)-monophosphatase 2, inositol(myo)-1(or 4)-monophosphatase 1, inositol(myo)-1(or 4)-monophosphatase 2 EST, Moderately similar to EF1G_HUMAN ELONGATION FACTOR 1-GAMMA [H.sapiens], ESTs, Highly similar to EF1G_HUMAN ELONGATION FACTOR 1-GAMMA [H.sapiens], Homo sapiens cDNA
167	4222	AA860024	a,bb		FLJ11216 fis, clone PLACE1008002, eukaryotic translation elongation factor 1 gamma

TABLES	S HUMAN	HONO COUTE AN	NOTATIONS .		7411 %. Docket No. 44921-5039W0 Doc. No. 1793397.1
56g. ID No.	ldentifer	GenBank Ace./ Ref. Seq. ID No.	Model Gode	Nama Homologous Gana	Homologous Cluster Namo
		Y.			EST, Moderately similar to 138369 beta tubulin [H.sapiens], EST, Weakly
					similar to 138369 beta-tubulin [H.sapiens], EST, Weakly similar to
					TUBULIN BETA-5 CHAIN [M.musculus], ESTs, Highly similar to
					A25113 tubulin beta chain 15 - rat [R.norvegicus], FK506-binding protein
					1A (12kD), RIKEN cDNA 2310061K05 gene, RIKEN cDNA 2410129E14
168	13974	AA860030	u,x,General		gene, tubulin, beta 2, tubulin, beta 5, tubulin, beta polypeptide
					Mus musculus 12 days embryo male wolffian duct includes surrounding
					region cDNA, RIKEN full-length enriched library, clone:6720466F14,
				hyaluronan mediated	full insert sequence, RIKEN cDNA 0610027D24 gene, TRAF4 associated
160	7000	A A 860030	,	motility receptor (RHAMM), hyaluronan-mediated motility receptor (RHAMM)	factor 1, hyaluronan mediated motility receptor (RHAMM), hyaluronan-mediated motility receptor (RHAMM)
169	7090	AA860039 AA860055	X	mounty receptor (KHAIMIM)	DKFZP547E2110 protein, hypothetical protein FLJ10604
170 171	23769 16323	AA866240	k,x w		protein PL3 10604
172	4462	AA866264	General		DEHYDROGENASE [R.norvegicus], Mus musculus 10 days embryo cDNA, RIKEN full-length enriched library, clone:2610528B18, full insert sequence, RIKEN cDNA 9430025F20 gene, Rattus norvegicus mRNA for 20- alpha-hydroxysteroid dehydrogenase (20-alpha-HSD), complete cds, aldo- keto reductase family 1, member C1, aldo-keto reductase family 1, member C4 (chlordecone reductase; 3-alpha hydroxysteroid dehydrogenase, type I; dihydrodiol dehydrogenase 4), expressed sequence Al315367, expressed sequence Al503553, hydroxysteroid (17-beta) dehydrogenase 5
					ESTs, Highly similar to A54602 microtubule-associated serine/threonine protein kinase MAST205 - mouse [M.musculus], ESTs, Moderately similar to A54602
					microtubule-associated serine/threonine protein kinase MAST205 - mouse [M.musculus], Homo sapiens cDNA: FLJ21699 fis,
					clone COL09829, KIAA0303 protein, KIAA0807 protein, Mus musculus adult male cecum cDNA, RIKEN full-length
					enriched library, clone:9130026D18, full insert sequence, microtubule associated testis specific
					serine/threonine protein kinase, syntrophin associated serine/threonine
173	15884	AA866276	k	I	kinase

TABLES	KAMUUH &	HOMOLOGUEAX	SHOUTATIONS :	enty t	Atty: Docket No. 44921-5039WO Doc. No. 1793397.1
Na	ldeniller	Consonk Aced Ref. Seq. ID No.	Model Gode	Homologous Cene Neme	Handlegous Cluster Name
174	17742	AA866302	c.y	4-hydroxyphenylpyruvate dioxygenase, 4- hydroxyphenylpyruvic acid dioxygenase	4-hydroxyphenylpyruvate dioxygenase, 4-hydroxyphenylpyruvic acid dioxygenase, ESTs, Weakly similar to HPPD MOUSE 4-HYDROXYPHENYLPYRUVATE DIOXYGENASE [M.musculus], ESTs, Weakly similar to S32820 alloantigen F-rat [R.norvegicus], hypothetical protein MGC15668
175	16333	AA866414	a,h	solute carrier family 4 (anion exchanger), member 1, solute carrier family 4, anion exchanger, member 1 (erythrocyte membrane protein band 3, Diego blood group)	ESTs, Highly similar to BAND 3 ANION EXCHANGE PROTEIN [M.musculus], ESTs, Weakly similar to B3HU band 3 anion transport protein, erythrocyte [H.sapiens], solute carrier family 4 (anion exchanger), member 1, solute carrier family 4, anion exchanger, member 1 (erythrocyte membrane protein band 3, Diego blood group)
176	18918	AA866444	p.q		
177 178	16853 18995	AA866454 AA866459	j,l,m,y,z h,m		
179 180 181	16013 26036 16059	AA866482 AA874849 AA874857	s r		ESTs, Highly similar to FGD1_HUMAN PUTATIVE RHO/RAC GUANINE NUCLEOTIDE EXCHANGE FACTOR [H.sapiens], ESTs, Weakly similar to FGD1 MOUSE PUTATIVE RHO/RAC GUANINE NUCLEOTIDE EXCHANGE FACTOR [M.musculus], ESTs, Weakly similar to FGD1_HUMAN PUTATIVE RHO/RAC GUANINE NUCLEOTIDE EXCHANGE FACTOR [H.sapiens], FGD1 family, member 3, RIKEN cDNA 5830461L01 gene, faciogenital dysplasia (Aarskog-Scott syndrome), faciogenital dysplasia homolog, faciogenital dysplasia homolog 2 (human)
182	16069 21633	AA874873 AA874951	f	, , , , , , , , , , , , , , , , , , , ,	ESTs, Weakly similar to RNA binding protein [H.sapiens]
184	16192	AA874995	w	· · · · · · · · · · · · · · · · · · ·	, , , , , , , , , , , , , , , , , , , ,
185	16254	AA875025	j		cellular retinoic acid-binding protein 1
187	20701	AA875032 AA875097	cc,General	fibrinogen, A alpha polypeptide	Homo sapiens clone HQ0582, expressed sequence Al303526, fibrinogen, A alpha polypeptide, fibrinogen, gamma polypeptide ADP-ribosylation factor 3, RIKEN
188	16416	AA875098	bb		cDNA 5430400P17 gene, Rattus norvegicus ADP-ribosylation factor 3 mRNA, complete cds, expressed sequence AA408731
189	16419	AA875102	bb		expressed sequence AL022645, expressed sequence C76690, small nuclear ribonucleoprotein E, small nuclear ribonucleoprotein polypeptide E
190	15313	AA875126	I,m,General		
191	10936	AA875146	W		
192	15371	AA875186 AA875205	u		ESTs, Weakly similar to IF39_HUMAN EUKARYOTIC TRANSLATION INITIATION FACTOR 3 SUBUNIT 9 [H.sapiens]

No.			NOTATIONS		Aliy, Docket No. 4/1921-5039/00 Doc. No. 1793397.1
104	ldeniller	ConBank Ace <i>l</i> Ref. Seq. ID No.:		Komologous Cene Name	Homologous Greter Name
194	15401	AA875257	x,z		
	15410	AA875268	p,s		NADH dehydrogenase (ubiquinone) Fe S protein 7 (20kD) (NADH-coenzyme Q reductase)
196	15420	AA875286	f		
197	15446	AA875327	s,w		
198	7936	AA875495	b,General		
199	17314	AA875509	i,l,m		EST, Moderately similar to A Chain A, Mdm2 Bound To The Transactivation Domain Of P53 {SUB 17-125 [H.sapiens], mouse double minute 2, human homolog of; p53-binding protein EST, Weakly similar to MOHU6N
	24472	AA875523	k		myosin alkali light chain 6, nonmuscle form [H.sapiens], myosin light chain, alkali, nonmuscle, myosin, light polypeptide 6, alkali, smooth muscle and non-muscle
201	15587	AA875577	j		
	15617	AA875620	General		
202	15618	AA875620	General		
					Jun-B oncogene, jun B proto-
203	5384	AA891041	f,cc,General		oncogene
	24814	AA891209	f,p		
205 2	21930	AA891322	d		nucleolin - rat [R.norvegicus], EST, Weakly similar to NUCL_HUMAN NUCLEOLIN [H.sapiens], ESTs, Highly similar to FUS_HUMAN RNA-BINDING PROTEIN FUS [H.sapiens], ESTs, Weakly similar to T17210 hypothetical protein DKFZp434N041.1 [H.sapiens], RIKEN cDNA 2700022N21 gene, fusion, derived from t(12;16) malignant liposarcoma, poly(A) binding protein, nuclear 1, small nuclear ribonucleoprotein 70kD polypeptide (RNP antigen) EST, Weakly similar to IF37 MOUSE EUKARYOTIC TRANSLATION INITIATION FACTOR 3 SUBUNIT 7 [M.musculus], eukaryotic translation initiation factor 3, subunit 7 (zeta,
					66/67 kDa), eukaryotic translation
206	1700F	A A 901552	h		initiation factor 3, subunit 7 (zeta,
	17225	AA891553	h		66/67kD)
	7522 9071	AA891571 AA891578	j,m b		ļ
					Homo sapiens, Similar to neurofilament, heavy polypeptide (200kD), clone MGC:20701 IMAGE:4634024, mRNA, complete cds, MAGE-E1 protein, MAGEF1 protein, Neurofilament, heavy polypeptide, RIKEN cDNA 2010107K23 gene, RIKEN cDNA 3830417A13 gene, general transcription factor IIH, polypeptide 1 (62kD subunit), melanoma antigen,
209	19321	AA891666	u	melanoma antigen, family D, 1	family D, 1, melanoma antigen, family D, 2, necdin, neurofilament, heavy polypeptide, neurofilament, heavy polypeptide (200kD)
	19321 17693	AA891666 AA891737	u j,l,m,n,y,z		D, 2, necdin, neurofilament, heavy polypeptide, neurofilament, heavy

TABLE S		HOWOLOGALE	NOTATIONS		Atty. Docket No. 44921-5089W0 Doc. No. 1793897.1
Seg. [D No.:	lden Maer	GenBenk Ace./ Ref. Seq. ID No.	Model Gode	Homologovs Gene Name	Homologous Cluster Name
213	18269	AA891769	General		ESTs, Highly similar to S03917 fibronectin ED-A [H.sapiens], ESTs, Moderately similar to Fourth And Fifth Fibronectin Type I Module Pair {SUB 183-275 [H.sapiens], Fibronectin 1, fibronectin 1
214	9905	AA891774	s,bb,General		
215	17061	AA891812	d		ESTs, Highly similar to 2013348A Ser kinase SRPK1 [H.sapiens], Mus musculus 13 days embryo head cDNA, RIKEN full-length enriched library, clone:3110005M20, full insert sequence, Mus musculus adult male lung cDNA, RIKEN full-length enriched library, clone:1200011B22, full insert sequence, SFRS protein kinase 1, SFRS protein kinase 2, serine/arginine-rich service proceific kinase 2
216	7050				rich protein specific kinase 2,
217	4463	AA891831	General		serine/threonine kinase 23 EST, Weakly similar to PE2R RAT 20- ALPHA-HYDROXYSTEROID DEHYDROGENASE [R.norvegicus], Mus musculus 10 days embryo cDNA, RIKEN full-length enriched library, clone:2610528B18, full insert sequence, RIKEN cDNA 9430025F20 gene, Rattus norvegicus mRNA for 20- alpha-hydroxysteroid dehydrogenase (20-alpha-HSD), complete cds, aldo- keto reductase family 1, member C1, aldo-keto reductase family 1, member C4 (chlordecone reductase; 3-alpha hydroxysteroid dehydrogenase, type I; dihydrodiol dehydrogenase 4), expressed sequence Al315367, expressed sequence Al503553, hydroxysteroid (17-beta) dehydrogenase 5
218	14289	AA891838	<u>i </u>		
219 220 221 222	17779 17438 22862	AA891842 AA891914 AA891943 AA891944	g,s,z General		EST, Weakly similar to ACY1_HUMAN AMINOACYLASE-1 [H.sapiens], aminoacylase 1
223	1159	AA891949	e,z	_	
224	4473	AA891965	General		
225	6362	AA892053	f,j,l,m		
226	9037	AA892066	у		EST, Weakly similar to PROD_HUMAN PROLINE OXIDASE, MITOCHONDRIAL PRECURSOR [H.sapiens], Homo sapiens mRNA for KIAA1653 protein, partial cds, proline dehydrogenase, proline oxidase), proline oxidase 1, proline oxidase
227	19469	AA892112	General		homolog
228	14595	AA892112 AA892128	o,t,v		nomolog
229	16527	AA892154	CC		
230	4482	AA892173	bb		
231	20917	AA892238	h		
232	2357	AA892268	d		Met proto-oncogene, RYK receptor-like tyrosine kinase, macrophage stimulating 1 receptor (c-met-related tyrosine kinase), met proto-oncogene

TABLE 8		HOMOLOGUE AN			Ally, Docker No. 44921-5039WG Doc. No. 1793397.1
Seg. (D No.	ldenlifter	Cenbank Ace./ Ref. Seq. ID No.	Model Gode	Homologous Gana Namo	Homologovs Cluster Name
233	18183	AA892271	h		
234	6523	AA892299	d		
					EST, Weakly similar to 60S RIBOSOMAL PROTEIN L3 [R.norvegicus], EST, Weakly similar to 184501 ribosomal protein L3 [H.sapiens], ESTs, Moderately similar to 60S RIBOSOMAL PROTEIN L3 [R.norvegicus], ESTs, Moderately similar to 184501 ribosomal protein L3 [H.sapiens], ESTs, Weakly similar to RL3 MOUSE 60S RIBOSOMAL PROTEIN L3 [M.musculus], RIKEN CDNA 1110057H16 gene, ribosomal
236	13647	AA892367	а		protein L3, ribosomal protein L3-like
237	17682	AA892378 AA892382	j,p,s,x,General	aldolase 2, B isoform,	ESTs, Weakly similar to T44342 hypothetical protein TSC501 [H.sapiens], kidney- and liver-specific gene, putative N-acetyltransferase Camello 2
				aldolase B, fructose-	
239	820	AA892395	g,s	bisphosphate	
240	14754	AA892414 AA892446	u		
241	17439	AA092446	<u> </u>		ubiquinol-cytochrome c reductase
242	16469	AA892462	р		(6.4kD) subunit
243	13609	AA892468 AA892468	i,General n,v,General		PROTEASE 7 PRECURSOR [M.musculus], ESTs, Weakly similar to MCT7 RAT MAST CELL PROTEASE 7 PRECURSOR [R.norvegicus], RIKEN cDNA 2410039E18 gene, RIKEN cDNA 4933401F05 gene, marapsin, mast cell protease 7, protease, serine, 21 (testisin), protease, serine, 22, protease, serine, 8 (prostasin), tryptase delta 1, tryptase, alpha EST, Weakly similar to MAST CELL PROTEASE 7 PRECURSOR [M.musculus], ESTs, Weakly similar to MCT7 RAT MAST CELL PROTEASE 7 PRECURSOR [R.norvegicus], RIKEN cDNA 2410039E18 gene, RIKEN cDNA 4933401F05 gene, marapsin, mast cell protease 7, protease, serine, 21 (testisin), protease, serine, 22, protease, serine, 8 (prostasin), tryptase delta 1, tryptase, alpha EST, Weakly similar to histone
					H2A.F/Z variant [H.sapiens], H2A histone family, member Z, RIKEN cDNA C530002L11 gene, histone
244	9254	AA892470	n,u		H2A.F/Z variant
245	11991	AA892483 	S		EST, Weakly similar to A32609 alpha- glucosidase [H.sapiens], ESTs, Weakly similar to LYAG MOUSE LYSOSOMAL ALPHA-GLUCOSIDASE PRECURSOR [M.musculus], alpha glucosidase 2, alpha neutral subunit,
246	1522	AA892486	f		glucosidase, alpha, acid ESTs, Highly similar to DS1_HUMAN DS-1 PROTEI [H.sapiens], immature
247	11994	AA892507	aa		colon carcinoma transcript 1
248	23888	AA892520	w		

FADIR	NAMES OF	HOMOLOGUE AN	SKICHTEATTONIS		** A11y. Docket No. 44921-5039W0
UABUE					Dog. No. 1798397.1
Seq. ID No.	ldenWiler	Consonk Ace <i>l</i> Ref. Seq. ID No.	Model Code	Komologovs Cene Name	Honologous Cluster Name
248	23889	AA892520	h		
249	8599	AA892522	р		
250	15154	AA802522			expressed sequence Al987846, expressed sequence AL023058, expressed sequence C77895, hypothetical protein MGC3178, protein disulfide isomerase-related protein, quiescin Q6
250	15154	AA892532 AA892545	p		quiesciii Qo
251 252	17468 11203	AA892554	f,h		· · · · · · · · · · · · · · · · · · ·
253	18906	AA892561	a,bb,General		
254	19327	AA892562	f.j.y,z		ESTs, Moderately similar to DKC1 RAT DYSKERIN [R.norvegicus], ESTs Weakly similar to DKC1_HUMAN DYSKERIN [H.sapiens], RIKEN cDNA 9030425C13 gene, dyskeratosis congenita 1, dyskerin, hypothetical protein, MGC:7014, nuclear factor of kappa light polypeptide gene enhancer in B-cells inhibitor-like 1
255	18274	AA892572	р		
256	4512	AA892578	cc		
257 258	15876 19085	AA892582 AA892598	w General		EST, Highly similar to 60S RIBOSOMAL PROTEIN L8 [R.norvegicus], EST, Moderately similar to 60S RIBOSOMAL PROTEIN L8 [R.norvegicus], EST, Weakly similar to JN0923 ribosomal protein L8, cytosolic [H.sapiens], ESTs, Highly similar to 60S RIBOSOMAL PROTEIN L8 [R.norvegicus], ESTs, Highly similar to RL8_HUMAN 60S RIBOSOMAL PROTEIN L [M.musculus], ESTs, Moderately similar to RL8_HUMAN 60S RIBOSOMAL PROTEIN L [M.musculus], expressed sequence AL024098, ribosomal protein L8
258	19086	AA892598	General		
259	20065	AA892647			EST, Highly similar to HISTONE H4 [R.norvegicus], ESTs, Highly similar to HISTONE H4 [R.norvegicus], H4 histone family, member D, H4 histone family, member D, H4 histone family, member H, H4 histone family, member K, Mus musculus 10 day old male pancreas cDNA, RIKEN full-length enriched library, clone:1810029H14, full insert sequence, Mus musculus 10 days embryo cDNA, RIKEN full-length enriched library, clone:2610027B07, full insert sequence, Mus musculus adult male testis cDNA, RIKEN full-length enriched library, clone:4930558J22, full insert sequence, Mus musculus adult male tongue cDNA, RIKEN full-length enriched library, clone:2310067E17, full insert sequence, histone 4 protein
260	20088	AA892666	a,n		
261	23783	AA892773	n		
	,	1 - 100-110	1	I	I

TABLES		KONOLOGALE W	INOTATIONS		Affy, Docket No. 44921-5039W0 Doc. No. 1793897.1
Seg. ID.	refilmed)	Consent Acel Ref. Seq. ID No.	Model Gode	Kamologous Gene Kamologous Gene	
262 263	17549 13542	AA892776 AA892798	f,z b		Mus musculus, Similar to solute carrier family 25 (mitochondrial carrier; phosphate carrier), member 3, clone MGC:7631, mRNA, complete cds, expressed sequence W51672, solute carrier family 25 (mitochondrial carrier; phosphate carrier), member 3
263	13542	AA892798	D		3-phosphoglycerate dehydrogenase,
					EST, Weakly similar to SERA RAT D-3 PHOSPHOGLYCERATE DEHYDROGENASE [R.norvegicus], ESTs, Moderately similar to SERA_HUMAN D-3-PHOSPHOGLYCERATE DEHYDROGENASE [H.sapiens], Mus musculus adult male testis cDNA, RIKEN full-length enriched library, clone:4930404C15, full insert sequence, RIKEN cDNA 1110059D05 gene, RIKEN cDNA 6430629L09 gene, glyoxylate reductase/hydroxypyruvate reductase, phosphoglycerate
264	22537	AA892799	General		dehydrogenase
					3-phosphoglycerate dehydrogenase, EST, Weakly similar to SERA RAT D-3 PHOSPHOGLYCERATE DEHYDROGENASE [R.norvegicus], ESTs, Moderately similar to SERA_HUMAN D-3-PHOSPHOGLYCERATE DEHYDROGENASE [H.sapiens], Mus musculus adult male testis cDNA, RIKEN full-length enriched library, clone:4930404C15, full insert sequence, RIKEN cDNA 1110059D05 gene, RIKEN cDNA 6430629L09 gene, glyoxylate reductase/hydroxypyruvate reductase, phosphoglycerate
264	22539	AA892799	V		dehydrogenase 3-phosphoglycerate dehydrogenase, EST, Weakly similar to SERA RAT D-3 PHOSPHOGLYCERATE DEHYDROGENASE [R.norvegicus], ESTs, Moderately similar to SERA_HUMAN D-3- PHOSPHOGLYCERATE DEHYDROGENASE [H.sapiens], Mus musculus adult male testis cDNA, RIKEN full-length enriched library, clone:4930404C15, full insert sequence, RIKEN cDNA 1110059D05 gene, RIKEN cDNA 6430629L09 gene, glyoxylate reductase/hydroxypyruvate reductase, phosphoglycerate
264	22538	AA892799	General		dehydrogenase
265	6951	AA892820	h		

TABLES	KAMOHI &	HOMOTOGAE AN	NOTATIONS		Aiiy. Docket No. 44921-5039WO Doc. No. 1793897.1
Seq. ID		GenBenk Acc./ Ref. Seq. ID No.	Model Gode	Homologous Cene ::; Name	I SECURITY CONTROLLER TO THE SECURITY OF THE S
		Joseph Marie Value of the Control of		19	ESTs, Highly similar to AR72_HUMAN
					AFLATOXIN B1 ALDEHYDE REDUCTASE 1 (AFB1-AR 1)
					(ALDOKETOREDUCTASE 7)
			i		[H.sapiens], ESTs, Moderately similar
					to AFAR RAT AFLATOXIN B1 ALDEHYDE REDUCTASE
					[R.norvegicus], RIKEN cDNA
					0610025K21 gene, aflatoxin B1
1					aldehyde reductase, aldo-keto
					reductase family 7, member A2 (aflatoxin aldehyde reductase), aldo-
					keto reductase family 7, member A3
266	23322	AA892821	j,z		(aflatoxin aldehyde reductase)
267	17923	AA892843	IT		RIKEN cDNA 2410042F05 gene,
					procollagen-lysine, 2-oxoglutarate 5-
					dioxygenase (lysine hydroxylase) 2,
					procollagen-lysine, 2-oxoglutarate 5- dioxygenase (lysine hydroxylase,
					Ehlers-Danios syndrome type VI),
					procollagen-lysine, 2-oxoglutarate 5-
268 269	22871 9053	AA892859	m 		dioxygenase 1
269	9053	AA892861	p,v,General		EST, Weakly similar to EFHU2
					translation elongation factor eEF-2
					[H.sapiens], ESTs, Highly similar to
					ELONGATION FACTOR 2 [R.norvegicus], U5 snRNP-specific
					protein, 116 kD, eukaryotic translation
					elongation factor 2, expressed
270	16482	AA892940	l _w		sequence Al451340, hypothetical protein FLJ21661
271	12020	AA893035	j,y		protein r Edz roo r
272	3863	AA893060	General		
273 274	13332 21305	AA893080 AA893082	i,General General		
275	16591	AA893191	j,z		
276	17447	AA893192	General		
277	3876	AA893205	n		Coloradulia 4 (abasabas dasa kisasa
					Calmodulin 1 (phosphorylase kinase, delta), Calmodulin III, ESTs, Highly
					similar to A Chain A, Calmodulin
					Complexed With Calmodulin-Binding
					Peptide From Smooth Muscle Myosin Light Chain Kinase (SUB 2-148
					[H.sapiens], RIKEN cDNA
					2310068O22 gene, calmodulin,
					calmodulin 1, calmodulin 1 (phosphorylase kinase, delta),
					calmodulin 2, calmodulin 2
					(phosphorylase kinase, delta),
278	3878	AA893230	General		calmodulin 3, calmodulin-like 3

TABLE 8	REMOVED IN	HOMOLOGUE AN	SKIOITATOKI		
					Dos. No. 1793997.
Seq. (i) (i) o.	ldenlifter	Cenbenk Aced Ref. Seq. ID No.	Model Code	ENES EVOCOLOMICAL	Homologous Cluster Name
279	20986	AA893242	q	fatty acid Coenzyme A ligase, long chain 2, fatty- acid-Coenzyme A ligase, long-chain 2	ESTs, Weakly similar to LCFB MOUSI LONG-CHAIN-FATTY-ACIDCOA LIGASE 2 [M.musculus], Mus musculus, Similar to fatty-acid-Coenzyme A ligase, long-chain 6, clone MGC:28744 IMAGE:4481949, mRNA, complete cds, Rattus norvegicus gonadotropin-regulated long chain acyl-CoA synthetase (GR-LACS) mRNA, complete cds, fatty acid Coenzyme A ligase, long chain 2, fatty-acid-Coenzyme A ligase, long-chain 1, fatty-acid-Coenzyme A ligase long-chain 2, lipidosin RIKEN cDNA 1300012C15 gene,
					RIKEN cDNA 2310076L09 gene, adipose differentiation related protein,
280	16168	AA893280	i,z,General		adipose differentiation-related protein
281	3886	AA893289	j,m,y		
282 283	15209 17800	AA893327 AA893436	y cc	· · · · · · · · · · · · · · · · · · ·	
284 285	17836 9084	AA893626 AA893717	h x		Guanine nucleotide-binding protein beta 1, Homo sapiens mRNA for FLJ00083 protein, partial cds, Mus musculus, clone MGC:7934 IMAGE:3583848, mRNA, complete cds, RIKEN cDNA 5930415H02 gene, WD repeat domain 5, guanine nucleotide binding protein (G protein), beta polypeptide 1, guanine nucleotide binding protein beta subunit 4, guanine nucleotide binding protein, beta 1, guanine nucleotide binding protein, beta 4, hypothetical protein, recombination protein REC14
286	22731	AA893743	d		
287	12031	AA893860	v		ESTs, Highly similar to YSHUT threoninetRNA ligase [H.sapiens], ESTs, Moderately similar to YSHUT threoninetRNA ligase [H.sapiens], hypothetical protein FLJ12528, threonyl-tRNA synthetase
288	17897	AA893905	k		
289	3447	AA893982	d		
290	22583	AA894009	n		
291	10540	AA894027	j		
292	4569	AA894059	x		

VABLE 3:	CHOMMON	HOMOLOGUEAX	NOTATIONS ::		Atty Docket No. 44921-5089W0
Seg. [D No.	logantifier	Conbenk Acel Ref. See. 10 No.	Model Gode #	Homológous Gene Namé	Doc: No. 1798397.1 Homologous Gluster Name
					Amyloid protein precursor-like protein 2, EST, Weakly similar to AMYLOID-LIKE PROTEIN 2 PRECURSOR [R.norvegicus], Human DNA sequence from clone RP3-461P17 on chromosome 20q12-13.2. Contains two novel genes, gene HE4 for Major Epididymis-specific protein E4 precursor (Epididymis Secretory protein E4), RPL5 (60S Ribosomal Protein L5), COX6C (Cytochrome C Oxidase subunit VIC) and HSPD1 (HSP60, Mitochondrial Matrix Protein P1 precursor, Heat Shock Protein 60, GROEL, HUCHA60) pseudogenes, the SPINT3 gene for Kunitz type serine protease inhibitor 3 (HKIB9), two genes for novel Kunitz/Bovine pancreatic trypsin inhibitor and WAP-type (Whey Acidic Protein) 'four-disulfide core' domains containing proteins and the gene for Eppin-1, -2 and -3. Contains ESTs, STSs, GSSs and a CpG island, RIKEN cDNA 1700024E17 gene, amyloid beta (A4) precursor-like protein 2, serine protease inhibitor, Kunitz type 2, tissue factor pathway inhibitor, tissue factor
	18419	AA894130	d		pathway inhibitor 2
	17336	AA894297	<u> </u>		
	19120	AA894318 AA899113	f,j		
296	19762	,			EST, Weakly similar to S45140 tubulin beta chain [H.sapiens], ESTs, Highly similar to T08726 tubulin beta chain [H.sapiens], ESTs, Highly similar to TBB1 RAT TUBULIN BETA CHAIN [R.norvegicus], ESTs, Moderately
298	18286 22051	AA899219 AA899498	u w	•	similar to I38370 beta-tubulin [H.sapiens], ESTs, Moderately similar to TBB1 RAT TUBULIN BETA CHAIN [R.norvegicus], RIKEN cDNA 2410129E14 gene, RIKEN cDNA 4930447K03 gene, RIKEN cDNA 4930542G03 gene, tubulin, beta 3, tubulin, beta polypeptide
298 : 298 :	22051 22052	AA899498 AA899498	q		similar to I38370 beta-tubulin [H.sapiens], ESTs, Moderately similar to TBB1 RAT TUBULIN BETA CHAIN [R.norvegicus], RIKEN cDNA 2410129E14 gene, RIKEN cDNA 4930447K03 gene, RIKEN cDNA 4930542G03 gene, tubulin, beta 3,
298 2 298 2 299 2	22051 22052 21628	AA899498 AA899498 AA899563			similar to I38370 beta-tubulin [H.sapiens], ESTs, Moderately similar to TBB1 RAT TUBULIN BETA CHAIN [R.norvegicus], RIKEN cDNA 2410129E14 gene, RIKEN cDNA 4930447K03 gene, RIKEN cDNA 4930542G03 gene, tubulin, beta 3,
298 2 298 2 299 3 300 4	22051 22052 21628 4262 4661	AA899498 AA899498 AA899563 AA899590 AA899709	q	receptor (calcitonin) activity modifying protein 3	similar to I38370 beta-tubulin [H.sapiens], ESTs, Moderately similar to TBB1 RAT TUBULIN BETA CHAIN [R.norvegicus], RIKEN cDNA 2410129E14 gene, RIKEN cDNA 4930447K03 gene, RIKEN cDNA 4930542G03 gene, tubulin, beta 3,
298 2 298 2 299 3 300 4	22051 22052 21628 4262	AA899498 AA899498 AA899563 AA899590	q aa i		similar to I38370 beta-tubulin [H.sapiens], ESTs, Moderately similar to TBB1 RAT TUBULIN BETA CHAIN [R.norvegicus], RIKEN cDNA 2410129E14 gene, RIKEN cDNA 4930447K03 gene, RIKEN cDNA 4930542G03 gene, tubulin, beta 3, tubulin, beta polypeptide receptor (calcitonin) activity modifying protein 3
298 298 299 300 301 302	22051 22052 21628 4262 4661	AA899498 AA899498 AA899563 AA899590 AA899709	q aa i t,General		similar to I38370 beta-tubulin [H.sapiens], ESTs, Moderately similar to TBB1 RAT TUBULIN BETA CHAIN [R.norvegicus], RIKEN cDNA 2410129E14 gene, RIKEN cDNA 4930447K03 gene, RIKEN cDNA 4930542G03 gene, tubulin, beta 3, tubulin, beta polypeptide receptor (calcitonin) activity modifying

				-201-	
		HOMOLOGUE AN	SKIOITATIONS		Atty: Docket No. 44921-5039WO Doc. No. 1798897.4
Seg. ID No	ldentifer	Consent Ace./ Rei. Sog. ID No.	Model Code	Homologous Gene Name	Homologous Cluster Name
305	23778	AA899854	c,k,x	topoisomerase (DNA) II alpha, topoisomerase (DNA) II alpha (170kD)	ESTs, Moderately similar to A40493 DNA topoisomerase [H.sapiens], ESTs, Moderately similar to TP2A MOUSE DNA TOPOISOMERASE II, ALPHA [M.musculus], topoisomerase (DNA) II alpha, topoisomerase (DNA) II beta, topoisomerase (DNA) II beta (180kD)
306 307	22060 9114	AA899898 AA899951	v,General		
308	8988	AA900148	f		
308	11841	AA900148	v		ESTs, Highly similar to IEFS_HUMAN TRANSFORMATION-SENSITIVE PROTEIN IEF SSP 3521 [H.sapiens], hypothetical protein FLJ12788
503	11071	I VIOUVET!			Alpha-2-macroglobulin, ESTs, Moderately similar to A2M2 MOUSE MURINOGLOBULIN 2 PRECURSOR [M.musculus], RIKEN cDNA 2610307I21 gene, alpha-2- macroglobulin, carbon catabolite repression 4 homolog (S. cerevisiae), expressed sequence AW456442,
310	4725	AA900290	cc		pregnancy-zone protein
311	4747	AA900465	General		
312	20988	AA900562	0		
313	3822	AA900863	b,g,General		DNA segment, Chr 17, human D6S81E 1, ESTs, Highly similar to S33681 translation initiation factor eIF-4A.I [H.sapiens], HLA-B associated transcript 1, Human clone 23933 mRNA sequence, eukaryotic translation initiation factor 4A, isoform 1, mitochondrial DEAD-box polypeptide 28, nuclear RNA helicase, DECD variant of DEAD box family
315	12420	AA901017	ь		Mus musculus, Similar to aspartyl- tRNA synthetase, clone MGC:6719 IMAGE:3586278, mRNA, complete cds, asparaginyl-tRNA synthetase, aspartyl-tRNA synthetase, hypothetical protein FLJ10514, hypothetical protein FLJ23441
316	4849	AA901155	S		
317	3959	AA901338	General	ļ	Consider Con line - ADD forming
318	22846	AA923982	a,d		succinate-CoA ligase, ADP-forming, beta subunit
319	4895	AA923999	k		
320	21546	AA924188	cc,General		
321	24192	AA924210	n,General		
322	4933	AA924301	g,I,General		ESTs, Weakly similar to NEUROFILAMENT TRIPLET H PROTEIN [M.musculus], RIKEN cDNA 0610009L18 gene, RIKEN cDNA 1300003A17 gene, RIKEN cDNA 2410142G14 gene, nucleolar protein
323	4944	AA924405	I,General		(KKE/D repeat)
324	4944	AA924403	r		(Care repeat)
J24	17340	/\\JZ4420	1'	I	<u> </u>

DEPENDENT PHOSPHATE TRANSPORT PROTEIN 2 Haap Homo sapiens, Similar to solute	TABLE 8		HOMOLOGAE AK	SMOTFATIONS		Atty, Docket No. 44 92 1-5039W0 Doc. No. 1798897.1
NPTZ_HUMAN RENALS SODIUM	899. ID No.	ldentifier	Coneank Aced Ref. Seq. ID No.	Model Code		Homologovis Cluster Name
3-phosphoglycerate dehydrogena SET, Weakly similar to SERA RA PHOSPHOGLYCERATE DEHYDROGENASE [R. norvegice ESTs, Moderately similar to SERA RA PHOSPHOGLYCERATE DEHYDROGENASE [R. norvegice ESTs, Moderately similar to SERA, HUMAN D-3- PHOSPHOGLYCERATE DEHYDROGENASE [H. sapiens], musculus adult male testis cDNA RIKEN full-length enriched library clone-4930404C15, full insert sequence, RIKEN DDNA 111005 gene, RIKEN DDNA 111005 gene, RIKEN DDNA 111005 gene, RIKEN DDNA 111005 gene, RIKEN DDNA 111005 gene, RIKEN DDNA 111005 gene, RIKEN DDNA 111005 gene, RIKEN DDNA 111005 gene, RIKEN CDNA 6430629L09 glyoxylate reductase/hydroxypri debudsae, phosphoglycerate dehydrogenase SET, Weakly similar to SERA RA PHOSPHOGLYCERATE DEHYDROGENASE [H. sapiens], musculus adult male testis cDNA RIKEN Lillength enriched library clone-4930404C15, full insert sequence, RIKEN cDNA 6430629L09 glyoxylate reductase/hydroxypri reductase, phosphoglycerate dehydrogenase dehydrogenase sequence, RIKEN cDNA 6430629L09 glyoxylate reductase/hydroxypri reductase, phosphoglycerate dehydrogenase sequence, RIKEN cDNA 6430629L09 glyoxylate reductase/hydroxypri reductase, phosphoglycerate dehydrogenase sequence, RIKEN cDNA 6430629L09 glyoxylate sequence, RIKEN cDNA 6430629L09 glyoxylate sequence, RIKEN cDNA 6430629L09 glyoxylate sequence, RIKEN cDNA 6430629L09 glyoxylate sequence, RIKEN cDNA 6430629L09 glyoxylate sequence, RIKEN cDNA 6430629L09 sequence, RIKEN cDNA 6430629L09 sequence, RIKEN cDNA 6430629L09 sequence, RIKEN cDNA 6430629L09 sequence, RIKEN cDNA 6430629L09 sequence, RIKEN cDNA 6430629L09 sequence, RIKEN cDNA 6430629L09 sequence, RIKEN cDNA 6430629L09 sequence, RIKEN cDNA 6430629L09 sequence, RIKEN cDNA 6430629L09 sequence, RIKEN cDNA 6430629L09 sequence, RIKEN cDNA 6430629L09 sequence, RIKEN cDNA 6430629L09 sequence, RIKEN cDNA 6430629L09 sequence, RIKEN cDNA 6430629L09 sequence, RIKEN cDNA 6430629L09 sequence, RIKEN cDNA	325	4949	AA924432	General		NPT2_HUMAN RENAL SODIUM-DEPENDENT PHOSPHATE TRANSPORT PROTEIN 2 [H.sapiens], Homo sapiens, Similar to solute carrier family 34 (sodium phosphate), member 1, clone MGC:18179 IMAGE:4155326, mRNA, complete cds, Rattus norvegicus mRNA for NaPi-2 alpha, complete cds, Solute carrier family 17 (sodium/hydrogen exchanger), member 2, expressed sequence Al649385, solute carrier family 34 (sodium phosphate), member 1, solute carrier family 34 (sodium phosphate),
3-phosphoglycerate dehydrogena SET, Weakly similar to SERA RA PHOSPHOGLYCERATE DEHYDROGENASE [R. norvegice ESTs, Moderately similar to SERA RA PHOSPHOGLYCERATE DEHYDROGENASE [R. norvegice ESTs, Moderately similar to SERA, HUMAN D-3- PHOSPHOGLYCERATE DEHYDROGENASE [H. sapiens], musculus adult male testis cDNA RIKEN full-length enriched library clone-4930404C15, full insert sequence, RIKEN DDNA 111005 gene, RIKEN DDNA 111005 gene, RIKEN DDNA 111005 gene, RIKEN DDNA 111005 gene, RIKEN DDNA 111005 gene, RIKEN DDNA 111005 gene, RIKEN DDNA 111005 gene, RIKEN DDNA 111005 gene, RIKEN CDNA 6430629L09 glyoxylate reductase/hydroxypri debudsae, phosphoglycerate dehydrogenase SET, Weakly similar to SERA RA PHOSPHOGLYCERATE DEHYDROGENASE [H. sapiens], musculus adult male testis cDNA RIKEN Lillength enriched library clone-4930404C15, full insert sequence, RIKEN cDNA 6430629L09 glyoxylate reductase/hydroxypri reductase, phosphoglycerate dehydrogenase dehydrogenase sequence, RIKEN cDNA 6430629L09 glyoxylate reductase/hydroxypri reductase, phosphoglycerate dehydrogenase sequence, RIKEN cDNA 6430629L09 glyoxylate reductase/hydroxypri reductase, phosphoglycerate dehydrogenase sequence, RIKEN cDNA 6430629L09 glyoxylate sequence, RIKEN cDNA 6430629L09 glyoxylate sequence, RIKEN cDNA 6430629L09 glyoxylate sequence, RIKEN cDNA 6430629L09 glyoxylate sequence, RIKEN cDNA 6430629L09 glyoxylate sequence, RIKEN cDNA 6430629L09 sequence, RIKEN cDNA 6430629L09 sequence, RIKEN cDNA 6430629L09 sequence, RIKEN cDNA 6430629L09 sequence, RIKEN cDNA 6430629L09 sequence, RIKEN cDNA 6430629L09 sequence, RIKEN cDNA 6430629L09 sequence, RIKEN cDNA 6430629L09 sequence, RIKEN cDNA 6430629L09 sequence, RIKEN cDNA 6430629L09 sequence, RIKEN cDNA 6430629L09 sequence, RIKEN cDNA 6430629L09 sequence, RIKEN cDNA 6430629L09 sequence, RIKEN cDNA 6430629L09 sequence, RIKEN cDNA 6430629L09 sequence, RIKEN cDNA 6430629L09 sequence, RIKEN cDNA	326	18891	AA924598	е		
327 22540 AA924630 V,General						DEHYDROGENASE [R.norvegicus], ESTs, Moderately similar to SERA_HUMAN D-3-PHOSPHOGLYCERATE DEHYDROGENASE [H.sapiens], Mus musculus adult male testis cDNA, RIKEN full-length enriched library, clone:4930404C15, full insert sequence, RIKEN cDNA 1110059D05 gene, RIKEN cDNA 6430629L09 gene, glyoxylate reductase/hydroxypyruvate
3-phosphoglycerate dehydrogena EST, Weakly similar to SERA RA PHOSPHOGLYCERATE DEHYDROGENASE [R.norvegice ESTs, Moderately similar to SERA, HUMAN D-3-PHOSPHOGLYCERATE DEHYDROGENASE [H.sapiens], musculus adult male testis cDNA RIKEN full-length enriched library clone:4930404C15, full insert sequence, RIKEN cDNA 111005; gene, RIKEN cDNA 6430629L09 glyoxylate reductase, phosphoglycerate dehydrogenase 4828 14759 AA92466 k 329 23123 AA924766 k 329 23123 AA924794 x 330 4067 AA924813 g.p. 43330 door AA924813 g.p. 5332 18130 AA924902 r.General 5332 18130 AA924904 d 5333 23141 AA925019 r ESTS, Weakly similar to A35863 tryptase [H.sapiens], ESTs, Weakly similar to MCT7 RAT MAST CELL PROTEASE 7 PRECURSOR [R.norvegicus], RIKEN cDNA 4733401093 gene, mast cell prof 7, tryptase delta 1, tryptase gamm tryptase, alpha 1335 21458 AA925049 f.aa,General tryptase, alpha 1335 21458 AA925049 f.aa,General 1536 5073 AA925061 m	327	22540	AA924630	v.General		
328	327	22541				3-phosphoglycerate dehydrogenase, EST, Weakly similar to SERA RAT D-3-PHOSPHOGLYCERATE DEHYDROGENASE [R.norvegicus], ESTs, Moderately similar to SERA HUMAN D-3-PHOSPHOGLYCERATE DEHYDROGENASE [H.sapiens], Musmusculus adult male testis cDNA, RIKEN full-length enriched library, clone:4930404C15, full insert sequence, RIKEN cDNA 1110059D05 gene, RIKEN cDNA 6430629L09 gene, glyoxylate reductase/hydroxypyruvate reductase, phosphoglycerate
330 4067 AA924813 g,p						, , ,
331 2888 AA924902 r,General	329	23123	AA924794	х		
331 2888 AA924902 r,General		4067				
333 23141 AA925019 r ESTs, Weakly similar to A35863 tryptase [H.sapiens], ESTs, Weakly similar to A35863 tryptase [H.sapiens], ESTs, Weakly similar to MCT7 RAT MAST CELI PROTEASE 7 PRECURSOR [R.norvegicus], RIKEN cDNA 4733401N09 gene, mast cell prof 7, tryptase delta 1, tryptase gammat cell prof 7, tryptase gammat cell prof 7, tryptase gammat cell prof 7, tryptase gammat cell prof 7, tryptase gammat cell prof 7, tryptase gammat cell prof 7, tryptase gammat cell prof 7, tryptase gammat cell prof 7, tryptase gammat cell prof 7, tryptase gammat cell prof 7, tryptase gammat cell prof 7, tryptase gammat cell prof 7, tryptase gammat cell prof 7, tryptase gammat cell prof 7, trypt	331	2888	AA924902	r,General		
333 23141 AA925019 r ESTs, Weakly similar to A35863 tryptase [H.sapiens], ESTs, Weakly similar to A35863 tryptase [H.sapiens], ESTs, Weakly similar to MCT7 RAT MAST CELI PROTEASE 7 PRECURSOR [R.norvegicus], RIKEN cDNA 4733401N09 gene, mast cell prof 7, tryptase delta 1, tryptase gammat ryptase, alpha 334 23195 AA925026 General tryptase, alpha 335 21458 AA925049 f,aa,General 336 5073 AA925061 m			AA924964			
ESTs, Weakly similar to A35863 tryptase [H.sapiens], ESTs, Weak similar to MCT7 RAT MAST CEL PROTEASE 7 PRECURSOR [R.norvegicus], RIKEN cDNA 4733401N09 gene, mast cell prof 7, tryptase delta 1, tryptase gamr tryptase, alpha 335 21458 AA925049 f,aa,General 336 5073 AA925061 m				r		
335 21458 AA925049 f,aa,General 336 5073 AA925061 m				General		tryptase [H.sapiens], ESTs, Weakly similar to MCT7 RAT MAST CELL PROTEASE 7 PRECURSOR [R.norvegicus], RIKEN cDNA 4733401N09 gene, mast cell protease 7, tryptase delta 1, tryptase gamma 1,
336 5073 AA925061 m						tryptase, alpha
337 14790 AA925087 o,General						

TABLE	BE CHUUMAN	HOMOLOGUE AN	NOTATIONS	and and	Atty, Docket No. 44921-5089W0
Seq. ID No:	ldentifier	Cenbenk Ace./ Ref. Seq. (D No.	Model Code	Homologous Gene : '- Name	Doc. No. 1793397.1 Homologous Gluster Name
	S Innovation of American				Homo sapiens, clone IMAGE:3940519, mRNA, partial cds, hypothetical protein
338	5089	AA925126	g		DKFZp762O076
					Mus musculus adult male testis cDNA, RIKEN full-length enriched library, clone:4930572N12, full insert
339	23261	AA925145	k,General		sequence, betaine-homocysteine methyltransferase, betaine- homocysteine methyltransferase 2
		1	1.,,		KIAA0438 gene product, Rattus
			1		norvegicus mRNA for
					neurodegeneration associated protein
					1, complete cds, expressed sequence AL022700, g1-related zinc finger
					protein, hypothetical protein,
					hypothetical protein FLJ11830 similar
					to Praja1, hypothetical protein, similar
340	17363	AA925150	а		to (U06944) PRAJA1, praja1
341	23448	AA925167	ļ!		
					EST, Weakly similar to TRI9_HUMAN THYROID RECEPTOR INTERACTING PROTEIN 9 [H.sapiens], ESTs, Highly
					similar to A44437 regenerating liver inhibitory factor RL/IF-1 - rat
					[R.norvegicus], ESTs, Weakly similar
					to I-kappa B alpha chain [M.musculus],
			Ì		nuclear factor of kappa light chain
					gene enhancer in B-cells inhibitor, alpha, nuclear factor of kappa light
					chain gene enhancer in B-cells
					inhibitor, beta, nuclear factor of kappa
					light polypeptide gene enhancer in B-
					cells inhibitor, alpha, nuclear factor of
					kappa light polypeptide gene enhancer
					in B-cells inhibitor, beta, nuclear factor
342	23159	AA925318	е		of kappa light polypeptide gene enhancer in B-cells inhibitor, epsilon
343	21500	AA925353	k		ermancer in B-cens inhibitor, epsilon
344	22479	AA925418	t		
345	21151	AA925539	b		
					EST, Moderately similar to 1604358A
					nuclear RNP protein L [H.sapiens],
					ESTs, Highly similar to 1604358A nuclear RNP protein L [H.sapiens],
					ESTs, Moderately similar to 1604358A
					nuclear RNP protein L [H.sapiens],
				heterogeneous nuclear	heterogeneous nuclear
346	16944	AA925541	f	ribonucleoprotein L	ribonucleoprotein L
			1		EST, Moderately similar to 1604358A
					nuclear RNP protein L [H.sapiens],
					ESTs, Highly similar to 1604358A nuclear RNP protein L [H.sapiens],
					ESTs, Moderately similar to 1604358A
					nuclear RNP protein L [H.sapiens],
				heterogeneous nuclear	heterogeneous nuclear
346	16945	AA925541	t	ribonucleoprotein L	ribonucleoprotein L
					ESTs, Moderately similar to JX0336
					succinate dehydrogenase [H.sapiens], succinate dehydrogenase complex,
347	17514	AA925554	bb		succinate denydrogenase complex, subunit A, flavoprotein (Fp)
348	5183	AA925662	i,General		oussillers, navoprotein (i p)
349	23189	AA925844	r		

TABLE	S THINIVIAN	HOMOLOGUE AN	PARONATOKI		Ally, Dodla, No. 44924-5089WG Dos. No. 1798897.1
899. [D No.	ldentifier	Ceneal Aced Ref. Seq. ID No.	Model (600)	Homologous Cene Name	Homologovs Cluster Name
					EST, Weakly similar to IMB3_HUMAN IMPORTIN BETA-3 SUBUNIT [H.sapiens], Homo sapiens cDNA FLJ12978 fis, clone NT2RP2006321, RAN binding protein 6, karyopherin
350	23190	AA925863	aa		(importin) beta 3
351	5252	AA926051	General		
352	22967	AA926080	h,cc		
353	17157	AA926129	b		
354	13411	AA926196	u,General		
355	5295	AA926247	General		potassium channel, subfamily K, member 1 (TWIK-1), potassium channel, subfamily K, member 3 (TASK-1), potassium channel, subfamily K, member 6 (TWIK-2), potassium channel, subfamily K, member 7
					DNA segment, human D4S114, P311
356	22928	AA926262	General		protein
357	8948	AA926316	r		
358	21798	AA926365	aa		CGI-69 protein, EST, Moderately similar to T43493 hypothetical protein DKFZp434C119.1 [H.sapiens], mitochondrial carrier family protein, mitochondrial solute carrier
359	9942	AA942697	s		Inna
					DNA segment, Chr 17, ERATO Doi
l		1			441, expressed, hematological and
360	6039	AA942716	x,General		neurological expressed 1
361	11174	AA942745	g,o,w		
362	23005	AA942770	9		
363 364	21318 6615	AA942774 AA942889	General v		
205	0004				RAT MACROPHAGE COLONY STIMULATING FACTOR I RECEPTOR PRECURSOR [R.norvegicus], colony stimulating factor 1 receptor, colony stimulating factor 1 receptor, formerly McDonough feline sarcoma viral (v-fms) oncogene
365	6691	AA943028	С		homolog
266	22442	AA042066			DEAD (aspartate-glutamate-alanine-aspartate) box polypeptide 5, DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 17 (72kD), KIAA0801 gene product, RIKEN cDNA 2610007K22 gene, RIKEN cDNA 4921506D17 gene, RIKEN cDNA 9130430L19 gene, RNA helicase, Rattus norvegicus RNA helicase with arginine-serine-rich domain mRNA, complete cds, expressed sequence Al325430, hypothetical protein, prp28, U5 snRNP 100 kd protein
366	22142	AA943066	р		100 kd protein
					ALEX1 protein, ALEX3 protein,
367	21993	AA943149	v,General		armadillo repeat protein ALEX2, hypothetical protein MGC3195
368	9061	AA943508	General		TU3A protein, hypothetical protein MGC11034
360	24390	ΔΔΩ/3531	hiny		ESTs, Weakly similar to VIL1 MOUSE VILLIN [M.musculus], actin binding LIM protein 1, advillin, erythrocyte membrane protein band 4.9 (dematin), and throcyte protein band 4.9 (villing protein band 4.9
369	24390	AA943531	b,j,n,y	L	erythrocyte protein band 4.9, villin

TABLES		HOMO FOGUE AV	SKOTATOKI		Atty. Docket No. 44924-5089WC
Seq. ID		Conbank Acci		Homologous Cone	Doc. No. 1793397.
ŊœĽ	Identifier	Ref. Seq. ID No.	Model Gode	Matima	Homologous Gluster Name
					EST, Moderately similar to I38369 beta tubulin [H.sapiens], EST, Weakly
					similar to 138369 beta-tubulin
					[H.sapiens], EST, Weakly similar to TUBULIN BETA-5 CHAIN
					[M.musculus], ESTs, Highly similar to
					A25113 tubulin beta chain 15 - rat
					[R.norvegicus], FK506-binding protein 1A (12kD), RIKEN cDNA 2310061K05 gene, RIKEN cDNA 2410129E14
					gene, tubulin, beta 2, tubulin, beta 5,
370	13976	AA943532	f,s,x		tubulin, beta polypeptide ajuba, expressed sequence Al481106,
371	22248	AA943537	cc,General		expressed sequence R75157, zyxin TATA box binding protein (TBP)-
0.70	00057	144040550	_		associated factor, RNA polymerase II,
372 373	22257 12673	AA943558 AA943773	m u.cc,General		J, 20kD
374	13641	AA944154	u		
375	2658	AA944155	f		
376	12770	AA944161	d		CDC28 protein bins - 2 DUCEN - DVA
	ļ				CDC28 protein kinase 2, RIKEN cDNA 1110038L14 gene, expressed
377	20903	AA944180	i,x		sequence AI047807
378	13507	AA944244	V		
379	15596 22681	AA944353	General		
380 381	6711	AA944413 AA944439	i,v,cc,General General		
					ESTs, Highly similar to AGP2_RAT ANGIOPOIETIN-2 (ANG-2) [R.norvegicus], angiopoietin 1, angiopoietin 2, angiopoietin-like 3,
382	14763	AA944481	i,q,General		angiopoietin-like 4, angiopoietin-like factor
383	22466	AA944605	h		
384 385	12301 7023	AA944727 AA944792	b d.m.aa		B-cell CLL/lymphoma 3, B-cell leukemia/lymphoma 3, ESTs, Highly similar to A44437 regenerating liver inhibitory factor RL/IF-1 - rat [R.norvegicus], ESTs, Weakly similar to I-kappa B alpha chain [M.musculus], molecule possessing ankyrin repeats induced by lipopolysaccharide (MAIL), homolog of mouse, nuclear factor of kappa light chain gene enhancer in B-cells 1, p105, nuclear factor of kappa light chain gene enhancer in B-cells inhibitor, alpha, nuclear factor of kappa light polypeptide gene enhancer in B-cells 2, p49/p100, nuclear factor of kappa light polypeptide gene enhancer in B-cells inhibitor, alpha, nuclear factor of kappa light polypeptide gene enhancer in B-cells inhibitor, alpha, nuclear factor of kappa light polypeptide gene enhancer in B-cells inhibitor, epsilon, testis-specific ankyrin motif containing protein
386	22536	AA944803	bb		
					CGI-89 protein, hypothetical protein DKFZp667O2416, hypothetical protein FLJ20984, leukocyte receptor cluster
387	22501	AA944811	g,l		(LRC) member 4
388	23967	AA944831	s		
389	26084	AA944922	<u> </u>		
390	11974	AA944958	General		l .

TABLE	BE CHUMAN	HOMOLOGUE AV	ROTATIONS		Atty/, Docket No. 44921-5089W0 Doc. No. 1799897.1
Seg. ID No. ***	CL CHARLESTON CO. T. T. T. T. T. T. T. T. T. T. T. T. T.	CenDenk Acc./ Ref. Seq. (D No.)	Model Code	Homologous Gene Name	Homologicus Aluster Mamo
392	22554	AA945076	z,General	The second secon	
393	14352	AA945181	General		
395	1798	AA945569	General		
396	22050	AA945604	i,aa		
397	19731	AA945615	d,o		
					Diaphorase (NADH/NADPH), NAD(P)H menadione oxidoreductase 2, dioxin inducible, NAD(P)H menadione oxidoreductase 2, dioxin-inducible, diaphorase (NADH/NADPH) (cytochrome b-5 reductase),
398	22612	AA945624	a,General		diaphorase 4 (NADH/NADPH)
399	22618	AA945656	aa		
400	11871	AA945679	V		
401	22656	AA945818	General		
402	6720	AA945828	р		
403	22351	AA945867	m		
404	22665	AA945877	f		
405	24243	AA945950	b		
406	22689	AA945962	General		
407	22692	AA945986	d	·	
408	22696	AA945996	c,General		
408	22697	AA945996	c,o		
409	22658	AA945998	w		
410	20832	AA946040	s		POLYPEPTIDE VIB [M.musculus], Human DNA sequence from clone RP4-591N18 on chromosome 22q13.1- 13.2 Contains a COX6B (Cytochrome C Oxidase subunit VIb (EC 1.9.3.1)) pseudogene, ESTs, GSSs and two putative CpG islands, RIKEN cDNA 2010000G05 gene, cytochrome c oxidase subunit VIb
411	18337	AA946046	General		
412	825	AA946108	General		EST, Highly similar to LMA3_HUMAN LAMININ ALPHA-3 CHAIN PRECURSOR [H.sapiens], ESTs, Highly similar to LMA3_HUMAN LAMININ ALPHA-3 CHAIN PRECURSOR [H.sapiens], Homo sapiens cDNA: FLJ21236 fis, clone COL01111, expressed sequence Al853660, laminin, alpha 3 (nicein (150kD), kalinin (165kD), BM600 (150kD), epilegrin)
413	8639	AA946221	e,cc,General		7
414	23237	AA946224	f		
415	15600	AA946250	o,aa		
416	19387	AA946275	t		actin related protein 2/3 complex, subunit 3 (21 kD)
417	6351	AA946344	d		EST, Weakly similar to JC5111 cyclindependent kinase-related protein 1b - rat [R.norvegicus], EST, Weakly similar to S10889 proline-rich protein [H.sapiens], ESTs, Highly similar to JC5111 cyclin-dependent kinase-related protein 1b - rat [R.norvegicus], Homo sapiens ALS2CR7 mRNA, complete cds, PCTAIRE protein kinase 1, PCTAIRE-motif protein kinase 1, PFTAIRE protein kinase 1
418	22057	AA946348	е		
	1				1
419 420	13962	AA946349 AA946351	aa		

TABLES	S HUMMAN	HOMOLOGUEAK	EKIDIVATIOKI		Atty: Docket No. 44921-5039W0 Doc: No. 1793397.1
Seq. ID No.	(Clemilifier	GenBenk Acc./ Ref. Seq. ID No.:	Model Gode	Hemelogous Gene Neme	Homologous Cluster Name
421	18280	AA946361	g	g ggddin administration in codd pack	EST, Moderately similar to 1923401A protein CBP [M.musculus], EST, Weakly similar to 1923401A protein CBP [M.musculus], ESTs, Highly similar to 1923401A protein CBP [M.musculus], ESTs, Weakly similar to 1923401A protein CBP [M.musculus], bromodomain, testis-specific, bromodomain-containing 2
422	18944	AA946391	v		
424	21410	AA946408	t		EST, Highly similar to HISTONE H4 [R.norvegicus], H4 histone family, member D, H4 histone family, member H, H4 histone family, member K, Mus musculus 10 day old male pancreas cDNA, RIKEN full-length enriched library, clone:1810029H14, full insert sequence, Mus musculus 10 days embryo cDNA, RIKEN full-length enriched library, clone:2610027B07, full insert sequence, Mus musculus adult male testis cDNA, RIKEN full-length enriched library, clone:4930558J22, full insert sequence, Mus musculus adult male tongue cDNA, RIKEN full-length enriched library, clone:2310067E17,
425	643	AA946439	о,у		full insert sequence, germinal histone H4 gene, histone 4 protein
426	20736	AA946443	×		EST, Weakly similar to NPD1_HUMAN NEURAL PROLIFERATION DIFFERENTIATION AND CONTROL PROTEIN-1 PRECURSOR (NPDC-1 PROTEIN) [H.sapiens], expressed sequence Al314472, neural proliferation, differentiation and control gene 1, neural proliferation, differentiation and control, 1
427	21878	AA946448	r		
428	21947	AA946451	bb		EST, Highly similar to AF151863 1 CGI 105 protein [H.sapiens]
429 430	17499 1809	AA946467	General x,General		
430	23360	AA946503 AA955104	f.General		
432	23471	AA955162	General		
433	9452	AA955206	b,General		
434	23512	AA955282	General		
434	22596	AA955298	General		
436	23283	AA955391	h		high density lipoprotein binding protein (vigilin)
437	23546	AA955393	General		\(\frac{1}{3\dots}\)
401	20070	, , , , , , , , , , , , , , , , , , , ,	Sinoral		ESTs, Weakly similar to SX10 RAT TRANSCRIPTION FACTOR SOX-10 [R.norvegicus], SRY (sex determining region Y)-box 10, SRY (sex determining region Y)-box 9 (campomelic dysplasia, autosomal sexreversal), SRY-box containing gene
438	12404	AA955408	b		10, expressed sequence AV220920
439	23626	AA955540	aa		

TABLES		HOMO FOOME W	SKOTATOKI		Auy, Docket No. 44921-5039W0 Doc No. 1793897.1
809. ID No.	ldenWier	Conbank Ace./ Ref. Seq. (D No.)		Homologous Gene Name	Homologous Clusier Namo
					EST, Highly similar to FBRL MOUSE FIBRILLARIN [M.musculus], EST, Weakly similar to A38712 fibrillarin [H.sapiens], ESTs, Highly similar to A38712 fibrillarin [H.sapiens], ESTs, Weakly similar to FBRL MOUSE FIBRILLARIN [M.musculus], expressed sequence AL022665,
441	17540	AA955914	bb		fibrillarin
442	24277	AA955962	General		
443	19939	AA955980	General		
					ESTs, Weakly similar to PCB3_MOUSE POLY(RC)-BINDING PROTEIN 3 (ALPHA-CP3) [M.musculus], ESTs, Weakly similar to ROK_HUMAN HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEIN K [R.norvegicus], IGF-II mRNA-binding protein 3, RIKEN cDNA 2610101N11 gene, coding region determinant-binding protein, heterogeneous nuclear
444	24000	AA956005			ribonucleoprotein K, poly(rC)-binding protein 3, poly(rC)-binding protein 4
445	11050	AA956164	s,v		EST, Weakly similar to T-COMPLEX PROTEIN 1, EPSILON SUBUNIT [M.musculus], ESTs, Moderately similar to T-COMPLEX PROTEIN 1, EPSILON SUBUNIT [M.musculus], T-complex 1, chaperonin containing TCP1, subunit 5 (epsilon), chaperonin subunit 5 (epsilon), expressed sequence AI528772, t-complex 1, t-complex protein 1
446	498	AA956278	a,General		Complex protein 1
447	23409	AA956294	q		
449	23773	AA956476	f,x		
450	23799	AA956530	d		
451 452	23800 23834	AA956534 AA956659	aa cc,General		ESTs, Weakly similar to RNG1_HUMAN RING1 PROTEIN [H.sapiens], ring finger protein 1, ring finger protein 2
702	23034	74,330003	cc,Ceneral		gene rich cluster, C8 gene,
453 454	16425 23847	AA956688 AA956723	f,x s		hypothetical protein MGC2577
455	23852	AA956746	j,l,m,z		ESTs, Weakly similar to CHD4_HUMAN CHROMODOMAIN HELICASE-DNA-BINDING PROTEIN 4 [H.sapiens], KIAA1416 protein, KIAA1696 protein, chromodomain helicase DNA binding protein 4
456	5989	AA956907	g,s		DnaJ (Hsp40) homolog, subfamily C, member 8, Homo sapiens mRNA; cDNA DKFZp434C2016 (from clone DKFZp434C2016), eukaryotic translation initiation factor 3, eukaryotic translation initiation factor 3, subunit 10 (theta, 150/170kD), expressed sequence C85189, guanylate kinase membraneassociated inverted 1, hypothetical protein DKFZp434B227, nasopharyngeal epithelium specific protein 1

TABLE	S: HIUMAN	HOMOLOGUE AV	MOTATIONS		Aliy, Docket No. 44921-5039W0 Doc. No. 1793897.1
Scg. ID No.	ldentifier	Cenbenk Ace./ Ref. Seq. ID No.		Homologove Gane Neme	Homologous Guster Name
					DnaJ (Hsp40) homolog, subfamily C, member 8, Homo sapiens mRNA; cDNA DKFZp434C2016 (from clone DKFZp434C2016), eukaryotic translation initiation factor 3, eukaryotic translation initiation factor 3, subunit 10 (theta, 150/170kD),
456	5990	AA956907	General		expressed sequence C85189, guanylate kinase membrane- associated inverted 1, hypothetical protein DKFZp434B227, nasopharyngeal epithelium specific protein 1
					X-linked protein, brain expressed, X- linked 1, hypothetical protein FLJ10097, nerve growth factor receptor (TNFRSF16) associated
457	23957	AA957123	u,General		protein 1 EST, Weakly similar to T12456 hypothetical protein DKFZp564M2423.1 [H.sapiens], ESTs, Highly similar to T12456 hypothetical protein DKFZp564M2423.1 [H.sapiens], PAI-1 mRNA-binding protein, intracellular hyaluronan-
458	22357	AA957264	General g,l,m,p,v,cc,Gene	-	binding protein
459	23314	AA957270	ral		
460 461	23995	AA957307	a,b General		EST, Moderately similar to G01026 serinetRNA ligase [H.sapiens], hypothetical protein FLJ20450, seryl- tRNA synthetase, uncharacterized gastric protein YC12P
462	24040	AA957422	c		CD3 antigen, zeta polypeptide, CD3Z antigen, zeta polypeptide (TiT3 complex), Fc fragment of IgE, high affinity I, receptor for; gamma polypeptide, Fc receptor, IgE, high affinity I, gamma polypeptide, expressed sequence Al573376
463	12478	AA957554	m		EST, Weakly similar to S01696 gene P3 protein [H.sapiens], ESTs, Highly similar to P3 PROTEIN [M.musculus], Protein P3
464	21306	AA957811	v		
465	24183	AA957889	t d		
466	17034	AA957905 AA963071	е		EST, Moderately similar to COPE_HUMAN COATOMER EPSILON SUBUNIT [H.sapiens], ESTs, Highly similar to COPE_HUMAN COATOMER EPSILON SUBUNIT [H.sapiens], coatomer protein complex, subunit epsilon, hypothetical protein FLJ13241
468	24053	AA963092	General		X-linked protein, brain expressed, X- linked 1, hypothetical protein FLJ10097, nerve growth factor receptor (TNFRSF16) associated protein 1
469	2767	AA963201	O		protein i
470	2022	AA963259	g		
471	2126	AA963488	đ		
472 473	24246 2195	AA963703 AA963746	b General		
	12100	17-10001 TO	Contra	İ	1

TABLE	OS CHOMMAN	HOMOLOGUEAN	SKIOITATIOKE		Atty. Docket No. 44921-5039000 Doc. No. 1793397.1
Seg. [D]	III AMA	ConBank Acci	Model Code	Homologous Cent; Name	Homologous Guster Name
No.	Identifier			Range	moniologous causter (kaine
475	2282	AA964147	е		
476	2284	AA964152	x		ubiquitin specific protease 14 (tRNA-
478	2350	AA964368	g,General		guanine transglycosylase) ESTs, Weakly similar to A29861 actin gamma [H.sapiens], ESTs, Weakly similar to 139393 alpha-actin [H.sapiens], ESTs, Weakly similar to S38782 actin beta chain [H.sapiens],
479	18830	AA964496	aa		Homo sapiens mRNA; cDNA DKFZp434B2115 (from clone DKFZp434B2115), RIKEN cDNA 1700052K15 gene, RIKEN cDNA 1700061J02 gene, actin-like 7a, expressed sequence AL023024, expressed sequence AV259599, melanoma X-actin, uncharacterized hypothalamus protein HARP11
480	2392	AA964541	b		
481	2395	AA964554	General		1
482	2410	AA964589	i,aa		
483	19145	AA964613	f	 	-
484	2424	AA964617	g		
485	3107	AA964687	General		
486	2457	AA964752	q,t		
487	6778	AA964763	b		
489	2468	AA964807	 		
403	2400	707304007	 		ESTs, Highly similar to GSH0 HUMAN
490 491	2469 12561	AA964814 AA964815	w General	glutamate-cysteine ligase , modifier subunit, glutamate cysteine ligase, modifier subunit	GLUTAMATECYSTEINE LIGASE REGULATORY SUBUNIT [H.sapiens], glutamate-cysteine ligase, modifier subunit, glutamate-cysteine ligase, modifier subunit
 	1.200				EST, Weakly similar to
492	2326	AA964892	laa		PROCOLLAGEN ALPHA 1(IV) CHAIN PRECURSOR [M.musculus], collagen, type IV, alpha 1, procollagen, type IV, alpha 3, procollagen, type IV, alpha 3, procollagen, type IV, alpha 5
493	21339	AA964962	General		ATP-binding cassette, sub-family A (ABC1), member 1, ATP-binding cassette, sub-family A (ABC1), member 12, ATP-binding cassette, sub-family A (ABC1), member 12, ATP-binding cassette, sub-family A (ABC1), member 7, EST, Moderately similar to ABC1_HUMAN ATP-BINDING CASSETTE, SUB-FAMILY A, MEMBER 1 [H.sapiens], EST, Weakly similar to ABC1 MOUSE ATP-BINDING CASSETTE, SUB-FAMILY A, MEMBER 1 [M.musculus], RIKEN cDNA 1810036E22 gene
494			General		KIKEN CDNA 1010030L22 gene
494	21390 12569	AA964988 AA965023	+		
+33	12303	AN300023	g	-	Homo sapiens, clone MGC:8857 IMAGE:3866266, mRNA, complete cds, inorganic pyrophosphatase,
496	2583	AA965166	bb		pyrophosphatase (inorganic) EST, Highly similar to T14795 hypothetical protein DKFZp434E171.1
497	15885	AA965207	r		[H.sapiens]
499	2905	AA996727	b,I,m,u,General		
500	2915	AA996782	u,bb		ESTs, Moderately similar to LAMIN B3 [M.musculus], hypothetical protein MGC2721, lamin B1, lamin B2
500					
500 501 502	2920 19525	AA996813 AA996856	d aa,General		

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TABLE 8	e Human	HOMOLOGUE AN	EKIÓIVATOĶI		Atty. Docket No. 44921-5089W0 Doc. No. 1798897.1
Scq. [D No.	ldenWier	ConBank Ace <i>l</i> Roi: Seq. ID No.	Model Gode	Homologous Cene Name	Homologous Gluster Name 🦠 -
503	2984	AA997015	С		
504	2986	AA997028	General		
505	3145	AA997237	General		
506	19249	AA997342	m		
507	16883	AA997345	General		
					methylene tetrahydrofolate dehydrogenase (NAD+ dependent), methenyltetrahydrofolate
508	3470	AA997374	p		cyclohydrolase, protease, serine, 15 EST, Weakly similar to LIS1 MOUSE PLATELET-ACTIVATING FACTOR ACETYLHYDROLASE IB ALPHA SUBUNIT [R.norvegicus], ESTs, Weakly similar to LIS1 MOUSE PLATELET-ACTIVATING FACTOR ACETYLHYDROLASE IB ALPHA SUBUNIT [R.norvegicus], WD repeat domain 3, f-box and WD-40 domain protein 2, hypothetical protein MGC2655, platelet-activating factor acetylhydrolase beta subunit (PAF-AH beta), platelet-activating factor acetylhydrolase, isoform 1b, beta1 subunit, platelet-activating factor acetylhydrolase, isoform Ib, alpha subunit (45kD)
510	3180	AA997425	<u> </u>	1	
511 512	3245 3020	AA997608 AA997656	General t		EST, Weakly similar to PLASMINOGEN ACTIVATOR INHIBITOR-2, TYPE A [R.norvegicus], expressed sequence Al876477, expressed sequence C76171, plasminogen activator inhibitor 2 type A, serine (or cysteine) proteinase inhibitor, clade B (ovalbumin), member 2, serine (or cysteine) proteinase inhibitor, clade B (ovalbumin), member 6, serine protease inhibitor 12
	-	7 4 400 7 400			Homo sapiens, clone IMAGE:4810400,
513 514	3269 3288	AA997800 AA997877	x,aa f		mRNA, antigen identified by monoclonal antibody Ki 67
515	23992	AA998164	k,x		DNA segment, Chr 4, ERATO Doi 639, expressed, EST, Moderately similar to CGB2 MOUSE G2/MITOTIC-SPECIFIC CYCLIN B2 [M.musculus], ESTs, Weakly similar to CGB1_HUMAN G2/MITOTIC-SPECIFIC CYCLIN B [H.sapiens], cyclin B1
					biliverdin reductase B (flavin reductase
516 517 518	3773 19623	AA998356 AA998422	General General		(NADPH)) B-cell CLL/lymphoma 3, molecule possessing ankyrin repeats induced by lipopolysaccharide (MAIL), homolog of mouse, nuclear factor of kappa light polypeptide gene enhancer in B-cells inhibitor, epsilon
519	3572	AA998516	x		ESTs, Weakly similar to G2/MITOTIC- SPECIFIC CYCLIN B1 [R.norvegicus], cyclin A2, cyclin B1, cyclin B1, related sequence 1, cyclin B2
520	2782	AA998565	C		cyclin-dependent kinase inhibitor 1C (P57), cyclin-dependent kinase inhibitor 1C (p57, Kip2)

TRABLE		HOMOFOGAE	EXIOTRATIONS		Atty, Docket No. 44921-5039We Doc. No. 17933976
Seq. [D No.	ldenillier	Conbank Aced Ref. Seq. ID No. AA998576	Model Code	Homologous Cene Name	Montologous Cluster Name
521	26119	AA998576	i,r,w,General		
522	22737	AA998660	aa		
523	3696	AA999030	е		
524	3079	AA999169	k,x,General		
				signal transducer and activator of transcription 1, signal transducer and activator of transcription 1,	Signal transducer and activator of transcription 3, expressed sequence AA408197, signal transducer and activator of transcription 1, signal transducer and activator of transcription 1, 91kD, signal transducer and activator of transcription 2, signal transducer and activator of transcription 3, signal transducer and activator of transcription 3 (acute-phase response factor), signal transducer and activator
525	3081	AA999171	e,p,r	91kD	of transcription 4
526	3082	AA999172	General	O IND	ESTs, Moderately similar to A54847 GMP synthase [H.sapiens], guanine monphosphate synthetase
527	17337	AB000717	k	_	Intemprioapriate synthetase
528	1535	AB000778	a		phospholipase D1, phophatidylcholine- specific, phospholipase D2
529	1382	AB002406	k	RuvB (E coli homolog)-like 1, RuvB-like protein 1	Homer, neuronal immediate early gene, 1B, Homer, neuronal immediate early gene, 2, homer, neuronal immediate early gene, 1, homer, neuronal immediate early gene, 2
530	20184	AB003753	d		
531	4312	AB010635	c,i,j,k,y,z		EST, Weakly similar to JC5408 carboxylesterase [H.sapiens], ESTs, Moderately similar to ES22 MOUSE LIVER CARBOXYLESTERASE 22 PRECURSOR [M.musculus], ESTs, Weakly similar to A48809 carboxylesterase [H.sapiens], ESTs, Weakly similar to JC5408 carboxylesterase [H.sapiens], T-complex expressed gene 5, carboxylesterase 1, carboxylesterase 2 (intestine, liver), carboxylesterase 3, carboxylesterase 3 (brain), hypothetical protein FLJ21736 DNA (cytosine-5-)-methyltransferase
532	21666	AB012214	k		1, DNA (cytosine-5-)- methyltransferase 2, DNA methyltransferase (cytosine-5) 1, EST Weakly similar to JE0378 DNA [R.norvegicus], Mus musculus DNA cytosine methyltransferase mRNA, f- box and leucine-rich repeat protein 11, protein containing CXXC domain 2 Mus musculus mouse-thyrotropin- releasing hormone receptor 2 (TRH- R2) mRNA, complete cds, thyrotropin
		į .			releasing hormone receptor, thyrotropin-releasing hormone receptor

TABLE 8	HUMAN	HOMOFOGALE (VI)			Atty. Docket No. 44921-5089W0
839. ID No.	ldenlijer	Cenbenk Ace <i>l</i> Ref. Seq. ID No		Homologous Gane Name	Homologous Cluster Name
<u>No.</u>	ldendifer.	Ref. Seq. ID No.+	Model Gode	<u>Name</u>	ESTs, Weakly similar to DUS8_HUMAN DUAL SPECIFICITY PROTEIN PHOSPHATASE 8 [H.sapiens], Human DNA sequence from clone RP11-243J16 on chromosome 20 Contains parts of 2 isoforms of the BCL2L1 (BCL2-like 1) gene, the gene for a novel protein (FLS353), the gene for a protein similar to MYLK (myosin, light polypeptide kinase), the FKHL18 (forkhead (Drosophila)-like 18) gene, part of three novel genes, ESTs, STSs, GSSs and CpG islands, KIAA1725 protein, dual specificity phosphatase 13, dual specificity
					phosphatase 14, dual specificity phosphatase 5, expressed sequence BB104621, expressed sequence
534	1183	AF013144	h		C79103, protein tyrosine phosphatase, non-receptor type 16 B-cell CLL/lymphoma 6, member B (zinc finger protein), ESTs, Moderately similar to zinc finger protein [H.sapiens], Homo sapiens, Similar to RIKEN cDNA 0610020102 gene, clone MGC:23427 IMAGE:4654320, mRNA,
535	1582	AF015911	h,z		complete cds
536	11483	AF020618	u,cc,General		HLA-G histocompatibility antigen,
537	20295	AF024712	aa		class I, G
538	19077	AF030358	ıy,z		hypothetical protein, clone 1-53, small inducible cytokine subfamily D (Cys-X3 Cys), member 1 (fractalkine, neurotactin), small inducible cytokine subfamily D, 1
539	23044	AF034218	General		RIKEN cDNA 4632428M18 gene, hyaluronidase 1, hyaluronidase 2, hyaluronoglucosaminidase 1, hyaluronoglucosaminidase 2, hyaluronoglucosaminidase 3, sperm adhesion molecule
540	25178	AF035955	d		ESTs, Moderately similar to hepatitis A
541	1564	AF035963	x,bb,General		virus cellular receptor 1 [H.sapiens], ESTs, Moderately similar to kidney injury molecule-1 [R.norvegicus], ESTs, Weakly similar to kidney injury molecule-1 [R.norvegicus]
542	8426	AF036335	f		ESTs, Moderately similar to NR54_HUMAN 54 KDA NUCLEAR RNA-BINDING PROTEIN [H.sapiens], ESTs, Moderately similar to p54nrb [H.sapiens], ESTs, Weakly similar to A54691 octamer-binding protein NonO - mouse [M.musculus], ESTs, Weakly similar to PSF_HUMAN PTB-ASSOCIATED SPLICING FACTOR [H.sapiens], RIKEN cDNA 5730470C09 gene, RIKEN cDNA 9030402K04 gene, non-POU-domain-containing, octamer-binding protein

TABLES	HUMAN	HOMOLOGUE AS	INOTATIONS		Ally. Docket No. 44921-5039WO Doc. No. 1793397.1
Seq. (D. No.	ldemiliter	ConBenk Acol Roi. Sog. ID No.	Model Gode	Homologous Gene Name	Homologous Cluster Namo
					ESTs, Highly similar to A55318 serine/threonine protein kinase [M.musculus], ESTs, Weakly similar to RIP MOUSE SERINE/THREONINE PROTEIN KINASE RIP [M.musculus], Human DNA sequence from clone RP5-1182A14 on chromosome 1 Contains part of a gene similar to rat Espin, a pseudogene similar to MST1 (macrophage stimulating 1 (hepatocyte growth factor-like)), a pseudogene similar to KIAA0454, a gene similar to MST1 (macrophage stimulating 1 (hepatocyte growth factor-like)), a pseudogene similar to KIAA0445, two isoforms of a novel gene (isoform 2 is the gene for KIAA1245 protein), ESTs, STSs, GSSs and CpG islands, ankyrin repeat domain 3, cerebral cavernous malformations 1, mitogen activated protein kinase kinase kinase 12, mitogen-activated protein kinase kinase 14, receptor (TNFRSF)-interacting serine-threonine kinase 1, receptor interacting protein 3, receptor
543	21817	AF036537	k	solute carrier family 1 (neuronal/epithelial high affinity glutamate transporter, system Xag),	interacting serine-threonine kinase 2
544	21145	AF038571	General	member 1, solute carrier family 1, member 1	
545	22602	AF044574	General		2,4-dienoyl CoA reductase 1, mitochondrial, 2,4-dienoyl CoA reductase 2, peroxisomal, 2-4-dienoyl- Coenzyme A reductase 2, peroxisomal, ESTs, Weakly similar to S11021 2,4-dienoyl-CoA reductase [R.norvegicus], Homo sapiens AS10 protein mRNA, partial cds, RIKEN cDNA 1200012F07 gene, RIKEN cDNA 2400003B18 gene, hydroxyprostaglandin dehydrogenase 15 (NAD), hydroxysteroid (17-beta) dehydrogenase 10, peroxisomal trans 2-enoyl CoA reductase; putative short chain alcohol dehydrogenase
546	13464	AF047707	h		UDP-glucose ceramide glucosyltransferase
547	12259	AF061266	h		EST, Highly similar to JC5807 trp3 protein - rat [R.norvegicus], EST, Weakly similar to TRP1_MOUSE TRANSIENT RECEPTOR POTENTIAL CHANNEL 1 (TRANSIENT RECEPTOR PROTEIN 1) (MTRP1) (TRP-RELATED PROTEIN 1) [M.musculus], transient receptor potential channel 1, transient receptor protein 1, transient receptor protein 3, transient receptor protein 4, transient receptor protein 5

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-2	13-

		HOMOLOGUE AX	SKONATOKI		Atty, Docket No. 44921-5089W0 Doc. No. 1793897.1
Seg. (D.) No.	ldentifier	ConBonk Ace./ Ref. Seq. ID No.	Model Code :-	Name : Homologous Gene	Emel refatib evogolomoth
					EST, Highly similar to A61209 hypertension-associated protein SA - rat [R.norvegicus], ESTs, Highly similar to A61209 hypertension- associated protein SA - rat [R.norvegicus], ESTs, Weakly similar to I54401 hypertension-associated protein SA [H.sapiens], KIAA1504 protein, SA (rat hypertension- associated) homolog, SA rat hypertension-associated homolog, expressed sequence AI788978, hypothetical protein FLJ20581, medium-chain acyl-CoA synthetase, solute carrier family 27 (fatty acid transporter), member 1, solute carrier family 27 (fatty acid transporter),
549	4589	AF062389	y,z		member 4 ESTs, Highly similar to 2008109A set
550	16007	AF062594	t	nucleosome assembly protein 1-like 1	gene [R.norvegicus], ESTs, Highly similar to SET_HUMAN SET PROTEIN [H.sapiens], SET translocation, SET translocation (myeloid leukemia-associated), nucleosome assembly protein 1-like 1, nucleosome assembly protein 1-like 2, nucleosome assembly protein 1-like 4 KIAA1348 protein, protein
					phosphatase 1G (formerly 2C), magnesium-dependent, gamma
551	15761	AF062741	u		isoform
552	17426	AF073839	p		
553	18615	AF074608	s		
554	15797	AF084205	f		ESTs, Moderately similar to T17365 serine/threonine protein kinase TAO1 - rat [R.norvegicus], ESTs, Weakly similar to ST25_MOUSE SERINE/THREONINE PROTEIN KINASE 25 (STERILE 20/OXIDANT STRESS-RESPONSE KINASE 1) (STE20/OXIDANT STRESS RESPONSE KINASE 1) (STE20-LIKE KINASE) [M.musculus], KIAA1361 protein, STE20-like kinase, expressed sequence AU020252, prostate derived STE20-like kinase PSK, serine/threonine kinase 10, thousand and one amino acid protein kinase
555	12932	AF102552	5		ESTs, Moderately similar to A55575 ankyrin 3, long splice form [H.sapiens], RIKEN cDNA 2310026G15 gene, RIKEN cDNA 2410004E01 gene, RIKEN cDNA 2410197A17 gene, RIKEN cDNA 2433400N19 gene, RIKEN cDNA 8430401K06 gene, RIKEN cDNA C430011H06 gene, ankyrin 3, node of Ranvier (ankyrin G), hypothetical protein FLJ20189, phospholipase A2, group VI, phospholipase A2, group VI (cytosolic, calcium-independent), proteasome (prosome, macropain) 26S subunit, non-ATPase, 10

TABLE		HOMOFOGALE VIX	INOTATIONS ?		Atty, Docket No. 44921-5039W0 Doc. No. 1793397.1
Seq. ID	i incompanya di di di di di di di di di di di di di	Centenik Ace.		Homologous Cene	
No.		Ref. Seq. ID No.	Model Gode	Name :	Homologous Cluster Name
556	18603	AI007649	×		EST, Moderately similar to A49013 tumor cell suppression protein HTS1 [H.sapiens], KIAA1277 protein, hypothetical protein FLJ22457, suppression of tumorigenicity 5
557	22733	AI007668	г		
558	22746	Al007672 .	r		
559	24109	AI007725	General		
					EST, Highly similar to HS9B RAT HEAT SHOCK PROTEIN HSP 90-BETA [R.norvegicus], EST, Weakly similar to HHMS84 heat shock protein 84 - mouse [M.musculus], ESTs, Highly similar to T46243 hypothetical protein DKFZp761K0511.1 [H.sapiens], expressed sequence AL022974, expressed sequence
500	45040	A1007820	<u></u>		C81438, heat shock 90kD protein 1,
561 562	10108 6804	AI007820 AI007857 AI007877	f General		beta, heat shock protein, 84 kDa 1 HGF-regulated tyrosine kinase substrate, Homo sapiens cDNA FLJ13428 fis, clone PLACE1002493, highly similar to Homo sapiens signal transducing adaptor molecule 2A (STAM2) mRNA, Mouse 31-kDa proline-rich salivary protein, complete cds of clone pUMP125, Mus musculus, Similar to proline-rich protein BstNI subfamily 2, clone MGC:18611 IMAGE:4165240, mRNA, complete cds, RIKEN cDNA 1700120F24 gene, RIKEN cDNA 4930406E12 gene, Rattus norvegicus proline-rich proteoglycan (PRPG2) mRNA, complete cds, proline-rich protein HaeIII subfamily 2, signal transducing adaptor molecule (SH3 domain and ITAM motif) 2
563 564	20099 11368	AI007893 AI007948	f,u d		
JU4	11300	NIOU 340			EST, Highly similar to HS9B RAT HEAT SHOCK PROTEIN HSP 90-BETA [R.norvegicus], EST, Weakly similar to HHMS84 heat shock protein 84 - mouse [M.musculus], ESTs, Highly similar to T46243 hypothetical protein DKFZp761K0511.1 [H.sapiens], expressed sequence AL022974, expressed sequence
565	15849	A1008074	h		C81438, heat shock 90kD protein 1, beta, heat shock protein, 84 kDa 1
566	3121	Al008160	General		CGI-83 protein

567 16	dentifier	Cerbank Acci Reil Seg. ID No.		Homologous Gene Name	EST, Highly similar to JC7290 guanine nucleotide binding protein G gamma 2 chain [H.sapiens], EST, Weakly similar to GBG9 RAT GUANINE NUCLEOTIDE-BINDING PROTEIN G(I)/G(S)/G(O) GAMMA-9 SUBUNIT [M.musculus], EST, Weakly similar to JC7290 guanine nucleotide binding protein G gamma 2 chain [H.sapiens], RIKEN cDNA 1110003P13 gene, guanine nucleotide binding protein (G protein), gamma 12, guanine nucleotide binding protein (G protein), gamma 2, guanine nucleotide binding protein (G protein), gamma 2 subunit, guanine nucleotide binding protein (G protein), gamma 3 subunit, guanine nucleotide binding protein (G protein), gamma 4 subunit, guanine nucleotide binding protein (G protein), gamma 4 subunit, guanine nucleotide binding protein (G protein), gamma 4 subunit, guanine nucleotide binding protein 4 EST, Moderately similar to CGB2 MOUSE G2/MITOTIC-SPECIFIC CYCLIN B2 [M.musculus], ESTs, Weakly similar to G2/MITOTIC-SPECIFIC CYCLIN B1 [R.norvegicus], Homo sapiens cDNA FLJ13342 fis,
		A1008190	t		nucleotide binding protein G gamma 2 chain [H.sapiens], EST, Weakly similar to GBG9 RAT GUANINE NUCLEOTIDE-BINDING PROTEIN G(I)/G(S)/G(O) GAMMA-9 SUBUNIT [M.musculus], EST, Weakly similar to JC7290 guanine nucleotide binding protein G gamma 2 chain [H.sapiens], RIKEN cDNA 1110003P13 gene, guanine nucleotide binding protein (G protein), gamma 12, guanine nucleotide binding protein (G protein), gamma 2, guanine nucleotide binding protein (G protein), gamma 2 subunit, guanine nucleotide binding protein (G protein), gamma 3 subunit, guanine nucleotide binding protein (G protein), gamma 4 subunit, guanine nucleotide binding protein 4 EST, Moderately similar to CGB2 MOUSE G2/MITOTIC-SPECIFIC CYCLIN B2 [M.musculus], ESTs, Weakly similar to G2/MITOTIC-SPECIFIC CYCLIN B1 [R.norvegicus],
		A1008190	t		nucleotide binding protein (G protein), gamma 4 subunit, guanine nucleotide binding protein 4 EST, Moderately similar to CGB2 MOUSE G2/MITOTIC-SPECIFIC CYCLIN B2 [M.musculus], ESTs, Weakly similar to G2/MITOTIC- SPECIFIC CYCLIN B1 [R.norvegicus],
		,			EST, Moderately similar to CGB2 MOUSE G2/MITOTIC-SPECIFIC CYCLIN B2 [M.musculus], ESTs, Weakly similar to G2/MITOTIC- SPECIFIC CYCLIN B1 [R.norvegicus],
568 12	2683		P		clone OVARC1001950, cyclin A1, cyclin B1, cyclin B1, related sequence
		AI008203	x		1, cyclin B2
569 22	2018	A1008309	Ь		ESTs, Highly similar to PROTO-ONCOGENE SERINE/THREONINE-PROTEIN KINASE PIM-1 [M.musculus], ESTs, Highly similar to S55333 protein kinase pim-2 [M.musculus], ESTs, Moderately similar to S55333 protein kinase pim-2 [M.musculus], Pim-1 oncogene, pim-1 oncogene, pim-2 oncogene, proviral integration site 1, serine threonine kinase pim3
		AI008309	n		phosphogluconate dehydrogenase
571 22	2599	AI008458	General		
		AI008578	p,General		
		AI008579 AI008629	r,x x		EST, Weakly similar to JH0446 75K autoantigen [H.sapiens], polymyositis/scleroderma autoantigen 1 (75kD)
575 38 576 39		A1008643	i,v,General		DnaJ (Hsp40) homolog, subfamily B, member 1, DnaJ (Hsp40) homolog, subfamily B, member 12, DnaJ (Hsp40) homolog, subfamily B, member 4, DnaJ (Hsp40) homolog, subfamily B, member 5, ESTs, Weakly similar to HS4L_HUMAN HEAT SHOCK 40 KDA PROTEIN 1 HOMOLOG [H.sapiens], RIKEN cDNA 1700029A20 gene, RIKEN cDNA 2010306G19 gene

TABLE	er human	HOMOROGAE VY	- SKOJTATOKI		Atty, Docket No. 44921-5089W0
509. [D] No.	(dentifier	Consonk Aced Ref. Seq. ID No.	Model Gode	Hemeleve Cene Neme	Homologous Gluster (Vaine)
577 578	7785 16701	A1008758 A1008838	aa q	dipeptidylpeptidase 4, dipeptidylpeptidase IV (CD26, adenosine deaminase complexing protein 2)	ESTs, Weakly similar to DPP4 MOUSE DIPEPTIDYL PEPTIDASE IV [M.musculus], ESTs, Weakly similar to DPP4 RAT DIPEPTIDYL PEPTIDASE IV [R.norvegicus], Homo sapiens chromosome 19, cosmid R26894, Homo sapiens, clone IMAGE:3447394, mRNA, partial cds, RIKEN cDNA 4932434F09 gene, dipeptidylpeptidase 4, dipeptidylpeptidase 6, dipeptidylpeptidase 8, fibroblast activation protein, fibroblast activation protein, alpha
579	21789	A1008930	k		EST, Moderately similar to CYSR RAT CYSTEINE-RICH PROTEIN 1 [R.norvegicus], ESTs, Weakly similar to S12658 cysteine-rich protein [H.sapiens], cysteine and glycine-rich protein 1, cysteine and glycine-rich protein 3 (cardiac LIM protein), cysteine rich protein, cysteine-rich protein 2, cysteine-rich protein 3, thymus LIM protein
580	21895	AI008971	General		
581	410	AI008974	i,aa,General		
582 583 584 585	21632 21596 22801 11876	Al009167 Al009168 Al009197 Al009321	General General General cc,General		BCL2-associated athanogene 2, ESTs, Highly similar to T08764 hypothetical protein DKFZp586C021.1 [H.sapiens]
586	2506	AI009341	General		
587	14370	A1009427	General k		EST, Weakly similar to PRCF_HUMAN PROTEASOME COMPONENT MECL-1 PRECURSOR [H.sapiens], ESTs, Weakly similar to PRCF_HUMAN PROTEASOME COMPONENT MECL-1 PRECURSOR [H.sapiens], proteasome (prosome, macropain) subunit, beta type 10, proteasome (prosome, macropain) subunit, beta type 7, proteasome (prosome, macropain) subunit, beta type 7, proteasome (prosome, macropain) subunit, beta type, 10, proteasome (prosome, macropain) subunit, beta type, 7
					ABP2_HUMAN ENDOTHELIAL ACTIN- BINDING PROTEIN [H.sapiens], ESTs, Moderately similar to
589 590	19275 4154	A1009460 A1009467	x g		ABP2_HUMAN ENDOTHELIAL ACTINBINDING PROTEIN [H.sapiens], filamin A, alpha (actin-binding protein-280), filamin B, beta (actin-binding protein-278)
590 591	4154 3464	AI009467 AI009589	g cc		BINDING PROTEIN [H.sapiens], filamin A, alpha (actin-binding protein- 280), filamin B, beta (actin-binding
590 591 592	4154 3464 3926	AI009467 AI009589 AI009592	g cc e		BINDING PROTEIN [H.sapiens], filamin A, alpha (actin-binding protein- 280), filamin B, beta (actin-binding
590 591	4154 3464	AI009467 AI009589	g cc		BINDING PROTEIN [H.sapiens], filamin A, alpha (actin-binding protein- 280), filamin B, beta (actin-binding

TABLES	K HUMAN	HOWO FORME W	A SHOTTATOLL		Atty. Docket No. 44921-5089W0
Seq. Id.		ConBenk Acc./		Homologous Ceine	<u>Doc. No. 1798397.1</u>
) Xo	ldentiller	Ref. Seq. ID No.	Model Goder :	Name:	Homologous Cluster Name
596	5458	Al009756	h	programmed cell death 6 interacting protein, programmed cell death 6- interacting protein	EST, Moderately similar to T14756 hypothetical protein DKFZp564F0923.1 [H.sapiens], EST, Weakly similar to A28996 proline-rich protein M14 precursor - mouse [M.musculus], EST, Weakly similar to PRP4_HUMAN SALIVARY PROLINE-RICH PROTEIN PO PRECURSOR [H.sapiens], expressed sequence AI462446, poly(A)-binding protein, nuclear 1, proline rich protein, proline rich protein 2, proline-rich protein BstNI subfamily 4, protein tyrosine phosphatase, non-receptor type 23
597	6844	AI009770	e,r,cc		EST Al317031, EST, Weakly similar to
598	15627	Al009810	aa		R3HU16 ribosomal protein S16, cytosolic [H.sapiens], expressed sequence AA420385, ribosomal protein S16
599 600	22619 7857	Al009825 Al009898	j,l,m,z		
601	13259	Al009898],1,111,2 r		
602	21105	AI010067	General		
603 604	24627 12716	AI010102 AI010178	aa General	testis enhanced gene transcript, testis enhanced gene transcript (BAX inhibitor 1)	CGI-119 protein, RIKEN cDNA 5031406P05 gene, testis enhanced gene transcript (BAX inhibitor 1) CGI-100 protein
605	18757	AI010216	aa		
606	2912	Al010220	aa,General		RIKEN cDNA 6720456116 gene, claudin 10, claudin 15, claudin 7
607	3316	AI010237	t		
608	15644	Al010256	General		ESTs, Highly similar to HISTONE H3.3 [R.norvegicus], H3 histone, family 3A, H3 histone, family 3B, H3 histone, family 3B (H3.3B)
609	657	Al010262	b		colony stimulating factor 2 receptor, beta 1, low-affinity (granulocyte- macrophage), interleukin 4 receptor, interleukin 4 receptor, alpha
610	3271	Al010303	b		
611	11081	Al010407	bb	ceruloplasmin,	DNA segment, Chr 3, ERATO Doi 555, expressed, EST, Highly similar to 1012298A factor VIIIC [H.sapiens], ESTs, Weakly similar to CERU MOUSE CERULOPLASMIN PRECURSOR [M.musculus], ESTs, Weakly similar to CERU RAT CERULOPLASMIN PRECURSOR [R.norvegicus], ESTs, Weakly similar to KUHU ferroxidase [H.sapiens], Hermansky-Pudlak syndrome 3, ceruloplasmin, ceruloplasmin
612	16521	AI010470	c,s,t,General	ceruloplasmin (ferroxidase)	(ferroxidase), hephaestin
613	6927	AI010542	General		
614 615	17524 6946	AI010568 AI010642	a,j,y,General n	growth hormone receptor	growth hormone receptor
					RIKEN cDNA 2510028H01 gene,
616	23509	AI010962	aa		sorting nexin 2, sorting nexin 3
617 618	6044 13855	AI011285 AI011361	t o		
619	21779	Al011380	cc		
	1	Al011460	cc		

TABLE	BE CHUMMAN	HOMOLOGALE VA	SKOLLÝLOKU	Company of the Compan	Aity. Docket No. 44921-5089W0 Doc. No. 1798897.1
Seg. [D] No.	Maniffler	Cenbenk Aced Ref. Seq. ID No.	Model Gode	Homologous Gane Name:	Homologous Chuster Namo
622	12629	Al011492	e,f		HYA22 protein, conserved gene amplified in osteosarcoma, nuclear LIM interactor-interacting factor
623	735	Al011560	f		Homo sapiens, Similar to RIKEN cDNA 2300002L21 gene, clone MGC:17528 IMAGE:3458906, mRNA, complete cds, RIKEN cDNA 2300002L21 gene, S100 calcium-binding protein A12 (calgranulin C), S100 calcium-binding protein, beta (neural), S100 protein, beta polypeptide, neural
					ESTs, Moderately similar to 2113291A laminin:SUBUNIT, Usher syndrome 2A (autosomal recessive, mild) homolog (human), hypothetical protein, MGC:
624	3941	Al011598	General		8159, laminin, alpha 5 butyrobetaine (gamma), 2-oxoglutarate dioxygenase (gamma-butyrobetaine hydroxylase) 1, epsilon-trimethyllysine
625	17550	Al011607	j,General		hydroxylase
626 627	10636 3995	Al011634 Al011678	e General		
628	16112	Al011706	h		EST, Weakly similar to SPLICING FACTOR, ARGININE/SERINE-RICH 5 [R.norvegicus], Mus musculus, Similar to splicing factor, arginine/serine-rich 7 (35kD), clone MGC:6268 IMAGE:2646366, mRNA, complete cds, neural-salient serine/arginine-rich, splicing factor, arginine/serine-rich 3 (SRp20), splicing factor, arginine/serine-rich 5, splicing factor, arginine/serine-rich 5 (SRp40, HRS) Fc fragment of IgG, high affinity Ia,
629	13354	Al011757	С		receptor for (CD64), Fc fragment of IgG, low affinity IIIa, receptor for (CD16), Fc fragment of IgG, low affinity IIIb, receptor for (CD16), Fc receptor, IgG, low affinity III, expressed sequence BB219290
630	12745	Al011799	СС	<u></u>	
631	18684	AI011812	t		DKFZP564O123 protein, putative breast adenocarcinoma marker (32kD)
632	4205	Al011982	b		
633	6518	AI012114	General		chromosome 1 open reading frame 25, hypothetical protein FLJ20244
634	17407	Al012145	General		
635	13093	AI012177	г		FK506 binding protein 4 (59 kDa), FK506 binding protein 8 (38 kDa), FK506-binding protein 4 (59kD), FK506-binding protein 6 (36kD), FK506-binding protein like, RIKEN cDNA 2210019E14 gene
636	15395	AI012216	f		Fas-associated factor 1, ORF, RIKEN cDNA 2210404D11 gene, UBX domain containing 2, expressed sequence AA408698, expressed sequence AI196514, putative glialblastoma cell differentiation-related

TABLE 3	RAMUH)	HOMOLOGUE AN	MOTATIONS		
6-06			[] [] [] [] [] [] [] [] [] []	[0]	<u>Doc. No. 1793397.1</u>
있으: Seed' (in)	ldentilier	ConBank Ace./ Ref. Seq. ID No.	Model Code	Homologous Caia Name	Honologous Cluster Name
637	21796	Al012221	d,General		EST X83352, ESTs, Highly similar to T17226 hypothetical protein DKFZp566G223.1 [H.sapiens], Homo sapiens, Similar to chloride intracellular channel 4, clone MGC:8812 IMAGE:3861372, mRNA, complete cds, RIKEN cDNA 5730531E12 gene, chloride intracellular channel 1, chloride intracellular channel 4, chloride intracellular channel 4 (mitochondrial), hypothetical protein DKFZp434N127, intracellular chloride ion channel protein p64H1
638 639	3981 6606	AI012235 AI012308	i,General i,r		
640	3417	Al012337	w		ESTs, Weakly similar to NHPX RAT NHP2/RS6 FAMILY PROTEIN YEL026W HOMOLOG [R.norvegicus], RIKEN cDNA 2410130M07 gene, non-histone chromosome protein 2 (S. cerevisiae)-like 1, nucleolar protein family A, member 2 (H/ACA small nucleolar RNPs), sperm specific antigen 1
641	24200	AI012356	b,t,General		andgen
642	7471	AI012379	cc		
643	7247	AI012438	9		
644	7127	Al012464	p,General		
645	3304	Al012471	b		
646	2311	AI012485	aa		
647	20817	AI012589	g,n,q		
648	3493	AI012590	v,General		
649	8975	AI012613	General		
650	11335	AI012619	ļ <u>. </u>		
651	21409	AI012637	General		
652 653 654	8476 4232	AI012638 AI012647 AI012958	w e,p,General		EST, Weakly similar to S33710 ribosomal protein S20, cytosolic [H.sapiens], Mus musculus, Similar to ribosomal protein S20, clone MGC:6876 IMAGE:2651405, mRNA, complete cds, expressed sequence AL024076
655	23128	Al013011	General		
656	20086	AI013260	General		
657	11969	Al013273	k		ESTs, Highly similar to A26061 gliaderived neurite promoting factor precursor [H.sapiens], ESTs, Highly similar to A27496 glia-derived nexin I alpha precursor [H.sapiens], ESTs, Weakly similar to GLIA DERIVED NEXIN PRECURSOR [R.norvegicus], serine (or cysteine) proteinase inhibitor, clade E (nexin, plasminogen activator inhibitor type 1), member 2, serine (or cysteine) proteinase inhibitor, clade I (neuroserpin), member 1
	26147	AI013387	aa		
658					T
659	8815	AI013437	p		
659 660	19722	AI013508	k		
659					

		HOMOLOGUE	EXPORTATION!		Atty. Docket No. 4/1921-5039WG Doc. No. 17/98897.1
800. ID 20.	ldentifier	CenBenk Ace./ Ref. Seq. ID No.	Model Code :	Homologous Gene Name	Honologovs Gveter Name
664	7274	Al013715	aa		Bone morphogenetic protein 6, bone morphogenetic protein 5, bone morphogenetic protein 6, bone morphogenetic protein 7, bone morphogenetic protein 7 (osteogenic protein 1), growth differentiation factor 15
665	7276	AI013730	е		
666	7278	AI013738	y,z,aa		ESTs, Weakly similar to S32567 A4 protein [H.sapiens], Homo sapiens, Similar to RIKEN cDNA 2900052H21 gene, clone MGC:21625 IMAGE:4214683, mRNA, complete cds, Homo sapiens, clone MGC:19762 IMAGE:3636045, mRNA, complete cds, proteolipid protein 2 (colonic
667	22592	AI013740	s,x,bb,General		epithelium-enriched) Mus musculus retinal cone arrestin 3 (Arr3) mRNA, complete cds, RIKEN cDNA 1200006117 gene, arrestin, beta 2, expressed sequence Al326910,
668	16584	AI013765	w		retinal S-antigen
669	24143	AI013804	j,i		
670 671	15928 21950	Al013829 Al013861	a,General j		3-hydroxyisobutyrate dehydrogenase, ESTs, Highly similar to D3HI_HUMAN 3-HYDROXYISOBUTYRATE DEHYDROGENASE, MITOCHONDRIAL PRECURSOR (HIBADH) [H.sapiens], cytokine-like nuclear factor n-pac
672	3260	AI013875	t		
673	2708	Al013882	d,q		
675 676	7299 15904	AI013886 AI013911 AI013971	p,r,t,General General		H.sapiens PABII pseudogene, Human DNA sequence from PAC 560B9 on chromosome 1q24-1q25. Contains profilin-like pseudogene, 60S ribosomal protein L4 pseudogene RNA binding protein, ESTs, GSS, Mus musculus adult male tongue cDNA, RIKEN full-length enriched library, clone:2310074E15, full insert sequence, RNA binding motif protein 3, RNA binding motif protein, X chromosome, RNA binding motif protein, X chromosome retrogene, cold inducible RNA-binding protein, testesspecific heterogenous nuclear ribonucleoprotein G-T
677	12781	AI014023	w		EST, Moderately similar to Y124_HUMAN HYPOTHETICAL PROTEIN KIAA0124 [H.sapiens], block of proliferation 1
				beta-carotene 15, 15'- dioxygenase, beta- carotene 15,15'-	EST, Moderately similar to 0806162D protein COII [M.musculus], EST, Weakly similar to 810024D cytochrome
678 679	19372 4241	AI014135	w	dioxygenase	oxidase II [H.sapiens] DKFZP564A2416 protein, EST, Moderately similar to T14738 hypothetical protein DKFZp564A2416.1 [H.sapiens], Homo sapiens cDNA FLJ14138 fis, clone MAMMA1002765, hypothetical protein FLJ13117

TABLES	HUMAN	HOMOTOGAE VI	MOTATIONS		: Ally, Docket No. 44921-5039WC
	E HISTORIA AND INCIDEN				Doc. No. 1798897.1
80g. ID No.	(dentifier	Conbank Ace <i>li</i> Rof. Soq. 1D No.	Model Gode	Homologous Cane : Name	Homologous Cluster Name
680	15247	Al014169	c,u		Homo sapiens cDNA: FLJ22783 fis, clone KAIA1993, Homo sapiens mRNA; cDNA DKFZp434B102 (from clone DKFZp434B102), KIAA1376 protein, expressed sequence AV216361, upregulated by 1,25-dihydroxyvitamin D-3 ESTs, Highly similar to JE0363
681	7315	AI028831	n		mitogen-activated protein kinase kinase kinase [H.sapiens], mitogen activated protein kinase kinase kinase 5, mitogen-activated protein kinase kinase kinase 6
682	16631	AI028856	General		
683	23297	AI028953	x		ESTs, Moderately similar to RUXG_HUMAN SMALL NUCLEAR RIBONUCLEOPROTEIN G [H.sapiens], small nuclear ribonucleoprotein polypeptide G
684	11326	Al029015	b		
685	2866	AI029058	n,y		
686	12812	Al029126	General		
687	17602	AI029156	р		
688	7392	AI029185	aa		
689 690	6517 7639	AI029264 AI029292	d,k,x		
					ESTs, Moderately similar to CB80_HUMAN 80 KDA NUCLEAR CAP BINDING PROTEIN [H.sapiens], Homo sapiens cDNA FLJ11599 fis, clone HEMBA1003879, nuclear cap
691	3874	AI029428	i,General		binding protein subunit 1, 80kD
692	12819 7452	AI029437 AI029466	r		
693 694	7452	AI029466 AI029608	b	-	
696	7537	AI029829	o,General		
697	2310	AI029969	V		
698	7585	AI030023	x		
699	7586	AI030024	b,n		
700	14492	AI030091	cc		
701	10673	Al030134	f		EST, Weakly similar to 1605244A erythrocyte ankyrin [H.sapiens], ESTs, Weakly similar to S68418 protein phosphatase 1M chain M110 isoform rat [R.norvegicus], Human DNA sequence from clone RP11-196N14 on chromosome 20 Contains ESTs, STSs, GSSs and CpG islands. Contains three novel genes, part of a gene for a novel protein similar to protein serine/threonine phosphatase 4 regulatory subunit 1 (PP4R1) and a gene for a novel protein with an ankyrin domain, RIKEN cDNA 1110058D09 gene, RIKEN cDNA 110058D09 gene, RIKEN cDNA 4930539L19 gene, expressed sequence AA408090, hypothetical protein MGC5540, leukocyte receptor cluster (LRC) member 3, myosin phosphatase, target subunit 1, protein phosphatase 1, regulatory (inhibitor) subunit 12A, testis-specific ankyrin motif containing protein
1702	1/615	IAI030163	lo.r		
702 703	7615 2370	AI030163 AI030179	o,r General		

TABLE	KINMMH] #	HOMOLOGUE A	NOTATIONS		Atty. Docket No. 44921-5039W0 Doc. No. 1793397.1
809. ID No. : : :	ldentifier	GenBenk Ace <i>l</i> Ref. Seq. ID No.	Model Gode :	Homologous Care	Homologous Cluster Name
705	7665	AI030472 AI030668	General t,bb	nucleosome assembly protein 1-like 1	ESTs, Highly similar to 2008109A set gene [R.norvegicus], ESTs, Highly similar to SET_HUMAN SET PROTEIN [H.sapiens], SET translocation, SET translocation (myeloid leukemia-associated), nucleosome assembly protein 1-like 1, nucleosome assembly protein 1-like 2, nucleosome assembly protein 1-like 4
707	24222	AI030704	k		
708	10740	AI030743	h		
709	10742	AI030773	е		
711 712	16169 19527	Al030932 Al030991	General f		RIKEN cDNA 1300012C15 gene, RIKEN cDNA 2310076L09 gene, adipose differentiation related protein, adipose differentiation-related protein
713	22614	AI031004	r		
714	3167	AI031012	e		ClpP (caseinolytic protease, ATP-dependent, proteolytic subunit, E. coli) homolog, caseinolytic protease, ATP-dependent, proteolytic subunit homolog (E. coli)
715	5350	AI043611	а		1
716	7858	AI043654	t	<u> </u>	
717	10784	AI043678	d		
718	9180	AI043694	aa		
719	7867	A1043695	aa		AMIDOPHOSPHORIBOSYLTRANSFE RASE PRECURSOR [H.sapiens], ESTs, Moderately similar to PUR1_HUMAN AMIDOPHOSPHORIBOSYLTRANSFE RASE PRECURSOR [H.sapiens], RIKEN cDNA 5730454C12 gene, expressed sequence AA675351, expressed sequence C79945, glutamine fructose-6-phosphate transaminase 2, glutamine-fructose-6-phosphate transaminase 2, phosphoribosyl pyrophosphate amidotransferase
720	7584	AI043724	General		
721	7895	AI043768	e		
722 723 724	7913 3899	AI043805 AI043849 AI043904	cc		ELL-RELATED RNA POLYMERASE II, ELONGATION FACTOR, ESTs, Highly similar to ELL2_HUMAN RNA POLYMERASE II ELONGATION FACTOR ELL2 [H.sapiens], ESTs, Weakly similar to ELL MOUSE RNA POLYMERASE II ELONGATION FACTOR ELL [M.musculus], ESTs, Weakly similar to ELL2_HUMAN RNA POLYMERASE II ELONGATION FACTOR ELL2 [H.sapiens], Mus musculus, clone IMAGE:3583970, mRNA, partial cds, Mus musculus, clone MGC:11987 IMAGE:3601737, mRNA, complete cds, eleven-nineteen lysine-rich leukemia gene, hypothetical protein FLJ22637
			I		
725	6766	AI043914	I Comment	"	
726	10818	AI043990	g,l,m,General		
727	7956	AI044018	f	l	

TABLE	B: HUMAN	HOMOFOGNE VI	SKIONTATIOKI		Atty, Doctot No. 44921-5039W0 Doc. No. 1793397.1
6- - 6		lossesses Assau	1		1.
Sed. שו No:	(dentifiter	ConBank Ace./ Rof. Seq. (D No.)	Model Code	Homologous Cene	Homologous Gluster Name
728	5393	AI044170	р		
729	5398	AI044177	q		
					EST, Weakly similar to S59856 collagen alpha 1(III) chain precursor - mouse [M.musculus], ESTs, Weakly similar to S59856 collagen alpha 1(III) chain precursor - mouse [M.musculus] Homo sapiens, Similar to hypothetical protein FLJ20783, clone MGC:1005 IMAGE:3139876, mRNA, complete cds, expressed sequence AW122071, hypothetical protein FLJ10355, procollagen, type XIX, alpha 1, sequence-specific single-stranded-
					DNA-binding protein, single-stranded
730	5425	AI044237	a.d		DNA-binding protein 2
731	8692	Al044247	r		2,4-dienoyl CoA reductase 1, mitochondrial, 2,4-dienoyl CoA reductase 2, peroxisomal, ESTs, Weakly similar to LUNG CARBONYL REDUCTASE [M.musculus], ESTs, Weakly similar to S11021 2,4-dienoyl-CoA reductase [R.norvegicus], FabG (beta-ketoacyl-[acyl-carrier-protein] reductase, E coli) like, Homo sapiens AS10 protein mRNA, partial cds, RIKEN cDNA 1200012F07 gene, RIKEN cDNA 1810027P18 gene, carbonyl reductase, carbonyl reductase 2, oxidoreductase UCPA
732	5430	AI044253	1		
733	5461	AI044338	g,p,General		
734	5464	AI044345	1		
735	3359	A1044347	aa		EST, Moderately similar to IL6B_HUMAN INTERLEUKIN-6 RECEPTOR BETA CHAIN PRECURSOR [H.sapiens], colony stimulating factor 3 receptor (granulocyte), cytokine receptor-like factor 1, interleukin 12 receptor, beta 2, interleukin 6 signal transducer, interleukin 6 signal transducer (gp130, oncostatin M receptor)
737 738	2695 5494	AI044396 AI044425	b General		oncostatin w receptor)
738 740	9882	A1044425	j,m		
741	5575	AI044688	g	 	
742	2348	AI044794	General		

TABLE &	(ALUMAN)	HOMOLOGUE AN	MOTATIONS		Ally, Dockel No. 44921-5189WO Doc. No. 1793897.1
Seq. ID No.	ldeniifier	Consink Aced Ref. Seq. ID No.	Model Gode	Homologove Gene Nemo	Homologove Gluster Name
					EST, Moderately similar to JH0148 nucleolin - rat [R.norvegicus], EST, Moderately similar to RBM8_HUMAN PUTATIVE RNA-BINDING PROTEIN 8 [H.sapiens], EST, Weakly similar to NUCL_HUMAN NUCLEOLIN [H.sapiens], ESTs, Highly similar to FUS_HUMAN RNA-BINDING PROTEIN FUS [H.sapiens], ESTs, Highly similar to RBM8_HUMAN PUTATIVE RNA-BINDING PROTEIN 8 [H.sapiens], ESTs, Moderately similar to RBM8_HUMAN PUTATIVE RNA-BINDING PROTEIN 8 [H.sapiens], Mus musculus pigpen protein mRNA, complete cds, RNA binding motif protein 8A, TATA box binding protein (TBP)-associated factor, RNA polymerase II, N, 68kD (RNA-binding protein 56), fusion, derived from
743	18205	AI044836	n		t(12;16) malignant liposarcoma
744	5626	AI044864	u		
745	5630	AI044869	f		
746	5634	AI044883	General		
747 748	4047 5654	AI044947 AI044976	l,m		
749	5684	AI044976	lv r		
750	19235	Al045074	General		ESTs, Highly similar to BGAL MOUSE BETA-GALACTOSIDASE PRECURSOR [M.musculus], ESTs, Weakly similar to BGAL MOUSE BETA GALACTOSIDASE PRECURSOR [M.musculus], Homo sapiens, clone IMAGE:3502329, mRNA, partial cds, Homo sapiens, clone IMAGE:3938286, mRNA, partial cds, RIKEN cDNA 4833408P15 gene, galactosidase, beta 1
751	5689	AI045074	i,aa,General		Deta 1
752	5711	AI045151	General		ESTs, Weakly similar to MCAT_HUMAN MITOCHONDRIAL CARNITINE/ACYLCARNITINE CARRIER PROTEIN [H.sapiens], expressed sequence AW108044, solute carrier family 25 (carnitine/acylcarnitine translocase), member 20, solute carrier family 25 (mitochondrial carnitine/acylcarnitine translocase), member 20, solute carrier family 25 (mitochondrial carrier; adenine nucleotide translocator), member 10, solute carrier family 25 (mitochondrial carrier; ornithine transporter), member 15, uncoupling protein 2 (mitochondrial, proton carrier), uncoupling protein 2, mitochondrial

Sept 19	TABLES	BE HOUMANN	HOMOLOGUE AV	SKOTIATOKI		Aity. Docket No. 44921-5089WO Doc. No. 1793897.1
ESTs, Modaretey similar to K882, MOUSE RIBOSOMAL PROTEIN 58 KINASE BETA 2 (58K-BETA 2) (70 KDA RIBOSOMAL PROTEIN 58 KINASE BETA 2 (58K-BETA 2) (70 KDA RIBOSOMAL PROTEIN 58 KINASE BETA 2 (58K-BETA 2) (70 KDA RIBOSOMAL SINASE BETA) (170 S6K	809. ID No.	ldeniiier			Hemologous Gene Neme:	
19237 Al045163 C C C C C C C C C						ESTs, Moderately similar to K6B2_MOUSE RIBOSOMAL PROTEIN S6 KINASE BETA 2 (S6K-BETA 2) (70 KDA RIBOSOMAL PROTEIN S6 KINASE 2) (P70-S6KB) (P70 RIBOSOMAL S6 KINASE BETA) (P70 S6KBETA) (S6K2) [M.musculus], NIMA (never in mitosis gene a)-related expressed kinase 3, RIKEN cDNA 2610318115 gene, expressed
19237 AI045163 C C C C C C C C C				:		
755 5735 A1045223 f 756 5474 A1045477 a. General 757 5811 A1045502 d.e 758 5819 A1045537 General 758 5819 A1045537 General 759 5839 A1045594 i 760 6808 A1045600 s 761 17755 A1045608 y 763 10020 A1045622 a 764 5855 A1045669 General 765 5881 A1045789 i 766 5897 A1045863 j 767 5890 A1045866 J 768 A1045866 J 769 A1045866 J 760 A104586	753	19237	AI045153	c		
1756 1757	754		AI045161	f		
TRAM-like protein, translocating chain associating membrane protein file associating membrane protein associating membrane protein file asso				f . O		
758 5819					-	<u> </u>
TRAM-like protein, translocating chain associaling membrane protein associaling membrane protein associaling membrane protein associaling membrane protein associaling membrane protein associaling membrane protein associaling membrane protein associaling membrane protein associaling membrane protein protein associaling membrane protein associaling membrane protein protein associaling membrane protein protein associaling membrane protein prot					 	
TRAM-like protein, translocating chain associating membrane protein				General	<u> </u>	
A	735	3039	A1045554	<u>'</u>	 	TRAM-like protein translocating chain-
761	760	6808	AI045600	s		
763 10020					 	associating memorane protein
Test					· · · · · · · · · · · · · · · · · · ·	· · · · · · · · · · · · · · · · · · ·
B aggressive lymphoma gene, DKFZP434J214 protein, KIAA1268 protein						
Protein FLJ13213, scaffold attachment factor B FLJ13213, scaffold attachment factor B	765	5881	AI045789	i		DKFZP434J214 protein, KIAA1268 protein
Total	700	5007	A10450C2	0		protein FLJ13213, scaffold attachment
EST, Weakly similar to C29149 proline rich protein - mouse [M.musculus], ESTs, Weakly similar to C29149 proline-rich protein - mouse [M.musculus], KIAA0999 protein, Mouse 31-kDa proline-rich salivary protein, complete cds of clone pUMP125, Mus musculus, Similar to proline-rich protein bstNI subfamily 2, clone MGC:18611 IMAGE:4165240, mRNA, complete cds, RIKEN cDNA 6030468819 gene, Rattus norvegicus proline-rich proteoglycan (PRPG2) mRNA, complete cds, proline-rich proteoglycan (PRPG2) mRNA, complete cds, proline-rich proteoglycan (PRPG2) mRNA, complete cds, proline-rich proteoglycan (PRPG2) mRNA complete cds, proline-rich protein Haelli subfamily 2 769 5329 Al045970 p 770 15093 Al058285 d 771 8002 Al058304 i 772 8017 Al058341 c Cdc42 effector protein 2, Cdc42 effector protein 2, Cdc42 effector protein 3 773 6828 Al058359 General 774 8177 Al058603 aa Cdc42 effector protein 3 775 3090 Al058730 aa 776 10093 Al058746 g 777 8143 Al058759 General 778 18659 Al058759 General 778 18659 Al05889 General 779 8163 Al05889 General 780 4789 Al05889 General 781 8221 Al059061 General					 	Tactor B
770	768	7540	Al045882	o,t,General		ESTs, Weakly similar to C29149 proline-rich protein - mouse [M.musculus], KIAA0999 protein, Mouse 31-kDa proline-rich salivary protein, complete cds of clone pUMP125, Mus musculus, Similar to proline-rich protein BstNI subfamily 2, clone MGC:18611 IMAGE:4165240, mRNA, complete cds, RIKEN cDNA 6030468B19 gene, Rattus norvegicus proline-rich proteoglycan (PRPG2) mRNA, complete cds, proline-rich
771 8002 Al058304 i 772 8017 Al058341 c 773 6828 Al058359 General effector protein 2, Cdc42 effector protein 3 774 8177 Al058603 aa 775 3090 Al058730 aa 776 10093 Al058746 g 777 8143 Al058759 General 778 18659 Al058762 f 779 8163 Al058837 aa 780 4789 Al058889 General 781 8221 Al059061 General						
772 8017 Al058341 c Cdc42 effector protein 2, Cdc42 effector protein 3 Cdc42 effector protein 3 effector protein 3 Al058359 General effector protein 3 aa Al058730 aa Al058730 aa Al058746 g Al058746 g Al058759 General Al058762 f Al058762 f Al0588762 f Al058837 aa Al058837 aa Al058837 Al058889 General Al058889 General Al058889 General Al058889 General Al058889 General Al058889 General Al058889 General Al058889 General Al058889 General Al058889 General Al058889 General Al058889 General Al058889 General Al058889 General General Al058889 General Al058889 General General Al058889 General Al058889 General General Al058889 General General General Al058889 General General Al058889 General General Al058889 General General General General Al058889 General Gen				i		
Cdc42 effector protein 2, Cdc42 effector protein 2, Cdc42 effector protein 3 774 8177 Al058603 aa 775 3090 Al058730 aa 776 10093 Al058746 g 777 8143 Al058759 General 778 18659 Al058762 f 779 8163 Al058876 General 780 4789 Al058889 General 781 8221 Al059061 General				c	<u> </u>	
774 8177 AI058603 aa 775 3090 AI058730 aa 776 10093 AI058746 g 777 8143 AI058759 General 778 18659 AI058762 f 779 8163 AI058837 aa 780 4789 AI058889 General 781 8221 AI059061 General						
775 3090 Al058730 aa 776 10093 Al058746 g 777 8143 Al058759 General 778 18659 Al058762 f 779 8163 Al058837 aa 780 4789 Al058889 General 781 8221 Al059061 General	774					
777 8143 Al058759 General 778 18659 Al058762 f 779 8163 Al058837 aa 780 4789 Al058889 General 781 8221 Al059061 General	775	3090	AI058730			
778 18659 AI058762 f 779 8163 AI058837 aa 780 4789 AI058889 General 781 8221 AI059061 General	776					
779 8163 AI058837 aa 780 4789 AI058889 General 781 8221 AI059061 General	777			General		
780 4789 AI058889 General 781 8221 AI059061 General				f		
781 8221 Al059061 General						
	781 782	10159	AI059061 AI059147	General d	ļ	

TABLES		HOMOFOGALE VA	INOTATIONS		
Seq. ID No.	ldeminer	GenBank Ace./ Ref. Seq. ID No.	Model Gode	Manielogous Cene Name	Honologous Gluster Namo
783	8245	Al059154	Ь		EST, Weakly similar to GBLP_HUMAN GUANINE NUCLEOTIDE-BINDING PROTEIN BETA SUBUNIT-LIKE PROTEIN 12.3 [H.sapiens], Homo sapiens mRNA for FLJ00083 protein, partial cds, IRA1 protein, Rattus norvegicus Sprague Dawley protein kinase C receptor mRNA, complete cds, WD repeat domain 5, expressed sequence AL033335, hypothetical protein, recombination protein REC14
784	8283	AI059290	n		
785	8314	AI059386	g,General		
786	10200	AI059444	li		
787	8347	AI059519	5		DKFZP566D213 protein, EST, Moderately similar to EPIDERMAL GROWTH FACTOR PRECURSOR [M.musculus], ESTs, Weakly similar to mel [M.musculus], Epidermal growth factor, epidermal growth factor, epidermal growth factor (beta- urogastrone), hypothetical protein MGC11256, nel-like 2 homolog (chicken)
788	18359	AI059675	n		EST, Highly similar to TERA HUMAN [H.sapiens], EST, Weakly similar to T46437 hypothetical protein DKFZp434K0126.1 [H.sapiens], ESTs, Weakly similar to T46437 hypothetical protein DKFZp434K0126.1 [H.sapiens], ESTs, Weakly similar to TERA HUMAN [H.sapiens], ESTs, Weakly similar to TERA RAT TRANSITIONAL ENDOPLASMIC RETICULUM ATPASE [R.norvegicus], RIKEN cDNA 4833413G10 gene, RIKEN cDNA 5430414H02 gene, spermatogenesis associated factor, valosin containing protein
789	10281	AI059947	b,t		S P P P P P P P P P P P P P P P P P P P
790	8494	AI059968	aa		
791 792	8495 8496	AI059971 AI059974	General General		Homo sapiens (clone NCD18) tumor necrosis factor receptor related protein mRNA, complete exon and repeat region, lymphotoxin B receptor, lymphotoxin beta receptor (TNFR superfamily, member 3), tumor necrosis factor receptor superfamily, member 8 KIAA1685 protein, KIAA1713 protein
793	10289	AI060053	i		CGI-142, RIKEN cDNA 3930401K13 gene
794	8548	AI060176	k		
795	8565	AI060236	t		
796	18322	AI060279	i,y,z		
797	8745 8785	AI070067	o		IK cytokine, down-regulator of HLA II, Mus musculus, Similar to IK cytokine, down-regulator of HLA II, clone MGC:25508 IMAGE:4920184, mRNA, complete cds
799	17506	A1070068	cc		growth arrest and DNA-damage- inducible 45 beta, growth arrest and DNA-damage-inducible, alpha, growth arrest and DNA-damage-inducible, beta

		HOWOROGUE A	EKOTATOKK	The state of the s	Atty, Docket No. 44921-5089Wo Doc. No. 1798897.1
(10) (10)	ldentifier	Cenbank Acel Ref. Seg. ID No.	Model Gode	Homologous Gene Kame	Honologous Cluster Name
800	9067	AI070087	General		ESTs, Highly similar to NUCL_HUMAN NUCLEOLIN [H.sapiens], Nucleolin, RIKEN cDNA 0610010A22 gene, eukaryotic translation initiation factor 3, subunit 4 (delta, 44 kDa), eukaryotic translation initiation factor 3, subunit 4 (delta, 44kD), nucleolin CGI-97 protein, EST, Weakly similar to YC97_HUMAN HYPOTHETICAL PROTEIN CGI-97 [H.sapiens], RIKEN cDNA 4733401P19 gene
802	4967	AI070179	k		glia maturation factor, gamma
803	18	AI070195	General		CGI-20 protein
804 805	24197 8869	Al070314 Al070330	General		armadillo repeat gene deletes in velocardiofacial syndrome, catenin (cadherin-associated protein), delta 1, catenin (cadherin-associated protein), delta 2 (neural plakophilin-related arm- repeat protein), plakophilin 4
806	8874	A1070336	h aa		·
807	10417	AI070336	b,cc m		<u> </u>
808	8901	AI070419	aa		toll-like receptor 1, toll-like receptor 10, toll-like receptor 2, toll-like receptor 6
809	14424	AI070421	I,p,General		ton into receptor 2, ten into receptor e
810	10434	AI070497	General		
811	8927	AI070523	v	· · · · · · · · · · · · · · · · · · ·	
812	8946	AI070611	q		
813	8950	AI070621	w		
814	8972	AI070673	General		
815	8981	AI070715	bb		
816 817 818	26184 3007 8999	A1070784 A1070824 A1070839	i.l w		ESTs, Highly similar to I49636 DNA-binding protein - mouse [M.musculus], ESTs, Moderately similar to I49636 DNA-binding protein - mouse [M.musculus], ESTs, Weakly similar to B57785 zinc finger protein ZNF136 [H.sapiens], ESTs, Weakly similar to OZF_HUMAN ZINC FINGER PROTEIN OZF [H.sapiens], Homo sapiens, Similar to zinc finger protein 136 (clone pHZ-20), clone MGC:10647 IMAGE:4053041, mRNA, complete cds, RIKEN cDNA 2310011F05 gene, pancreas zinc finger protein, zinc finger protein 136 (clone pHZ-20), zinc finger protein 260, zinc finger protein 63, zinc finger protein 97 DKFZP564F0522 protein, ESTs, Weakly similar to T08675 hypothetical protein DKFZp564F0522.1 [H.sapiens]
 	10000	7.07003	P		ESTs, Highly similar to NRP2_RAT
819 820 821	10477 24301 8721	Al070868 Al070911 Al071024	e,f k General		NEUROPILIN-2 PRECURSOR (VASCULAR ENDOTHELIAL CELL GROWTH FACTOR 165 RECEPTOR 2) [R.norvegicus], neuropilin 2, neuropilin-2, platelet derived growth factor C
822	9212	A1071098	х		
823	1831	Al071137	С		cell division cycle 25 homolog B (S. cerevisiae), cell division cycle 25 homolog C (S. cerevisiae), cell division cycle 25B, cell division cycle 25C, expressed sequence Al604853

TABLES	RAMUMAN	HOMOLOGUE AN	SKIOTKATIOKI		Allys Docket No. 44921-5039W0 Docs No. 1793397.1
	ldentifier	ConBenk Ace./ Ref. Seq. (D No.	Model Gode	Homologous Gene Nama	Homologous Gluster Name
824	11005	AI071139	r	Commence and the second process of the second	
GET	11000	7.40,7100			ESTs, Highly similar to ROG_HUMAN HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEIN G [H.sapiens], Homo sapiens, Similar to RNA binding motif protein, X chromosome, clone MGC:9398 IMAGE:3875565, mRNA, complete cds, RIKEN cDNA 1700012H05 gene, RNA binding motif protein, X chromosome, RNA binding motif protein, X chromosome retrogene, RNA binding motif protein, Y chromosome, family 1, member A1, testes-specific heterogenous nuclear
825	9104	AI071173	j,m		ribonucleoprotein G-T
826	9583	AI071185	General		
827	9644	AI071410	С		
920	16050	A1071400	Conoral		serine palmitoyltransferase, long chain base subunit 2
828 829	16058 11057	AI071490 AI071509	General f,o		Dase Suburiil Z
831	5695	Al071566	bb		
832	9671	AI071568	w		
032	3071	7107 1000	 ''		DNA segment, human D4S114, P311
833	22929	AI071578	General		protein
834	9673	AI071581	General		
835	9699	AI071646	General		
837	9799	AI072008	q,y,z		
838	9808	AI072050	d		
839	22796	AI072213	General		
840	9271	A1072405	v		
841	10869	AI072425	w		
842	21797	A1072439	General		EST X83352, ESTs, Highly similar to T17226 hypothetical protein DKFZp566G223.1 [H.sapiens], Homo sapiens, Similar to chloride intracellular channel 4, clone MGC:8812 IMAGE:3861372, mRNA, complete cds, RIKEN cDNA 5730531E12 gene, chloride intracellular channel 1, chloride intracellular channel 4, chloride intracellular channel 4 (mitochondrial), hypothetical protein DKFZp434N127, intracellular chloride ion channel protein p64H1
843	9306	AI072521	r		protein pour r
844	9312	AI072550	 		
845	10893	AI072559	x	l	
846	1501	AI072634	cc,General		
847	6548	AI072658	General		
848	9363	AI072695	d		DnaJ (Hsp40) homolog, subfamily C, member 4, Homo sapiens, clone MGC:19482 IMAGE:4309314, mRNA, complete cds, hypothetical protein FLJ11506
850	9409	AI072841	n		
851	9410	AI072842	w		
852	9468	AI073021	General		
853	9518	AI073223	f		
					EST, Moderately similar to CARBONIC ANHYDRASE II [R.norvegicus], carbonic anhydrase 2, carbonic anhydrase VIII, carbonic anhydrase VIII,
854	11183	Al100768	<u> t </u>		carbonic anhydrase-like sequence 1
855	9190	AI100835	е	<u> </u>	

TABLES	REWINDER &	HOMOFOGALE VI	EKIO TRATI ONI		. Aiiy. Dockei No. 44921-5039WO Doc. No. 1793397.1
809. ID No. +	(dentifier	ConBank Ace <i>l</i> Ref. Seq. ID No.	Model Gode	Homologovs Gene Neme	Honologous Givster Name
856	2029	AI100842	р		
857	5687	AI101006	e		
858	15192	Al101099	g,cc		ESTs, Moderately similar to AF078844 1 hqp0376 protein [H.sapiens], expressed sequence AA409533 ATP synthase, H+ transporting,
859	17399	Al101157	o		mitochondrial F0 complex, subunit f, mitochondrial F0 complex, subunit f, isoform 2, EST, Weakly similar to ATPK_HUMAN ATP SYNTHASE F CHAIN, MITOCHONDRIAL [H.sapiens], ESTs, Highly similar to ATPK_HUMAN ATP SYNTHASE F CHAIN, MITOCHONDRIAL [H.sapiens]
860	9339	Al101160	l,m,o		
					ESTs, Moderately similar to HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEIN C [R.norvegicus], Mus musculus highglycine/tyrosine protein type I E5 mRNA, complete cds, RNA binding protein p45AUF1, expressed sequence C85084, heterogeneous nuclear ribonucleoprotein A/B, heterogeneous nuclear ribonucleoprotein D, heterogeneous nuclear ribonucleoprotein D (AU-rich element RNA-binding protein 1, 37kD),
l			١		heterogeneous nuclear
861	6321	Al101256	General		ribonucleoprotein D-like
862	5421	Al101270	c		Rho GDP dissociation inhibitor (GDI) beta, expressed sequence C87222, rho, GDP dissociation inhibitor (GDI) beta
		A1101270			ets variant gene 5 (ets-related
863	11910	AI101323	General		molecule)
864	23140	AI101608	e		
865	4119	AI101901	General		
866	16324	Al102009	b		
867	18642	AI102023	o		brain-specific membrane-anchored protein, chromosome 1 open reading frame 8
868	19373	Al102044	а	beta-carotene 15, 15'- dioxygenase, beta- carotene 15,15'- dioxygenase, frizzled (Drosophila) homolog 1, frizzled homolog 1, (Drosophila)	EST, Moderately similar to 0806162D protein COII [M.musculus], EST, Weakly similar to 810024D cytochrome oxidase II [H.sapiens]
869	7051	Al102055	h	,	ESTS, Highly similar to 2013348A Ser kinase SRPK1 [H.sapiens], Mus musculus 13 days embryo head cDNA, RIKEN full-length enriched library, clone:3110005M20, full insert sequence, Mus musculus adult male lung cDNA, RIKEN full-length enriched library, clone:1200011B22, full insert sequence, SFRS protein kinase 1, SFRS protein kinase 2, serine/argininerich protein specific kinase 2, serine/threonine kinase 23
	6544	AI102055	in ic		Sering/uncomine killase 23
870 871			w		
871	10227 23849	AI102248 AI102318	· · · · · · · · · · · · · · · · · · ·		
873	11954	AI102505	le,q g,j,s		
 , , , , , , , , , , , , , , , , , , ,	1.1007		٠٠،٠٠	·	TYRO protein tyrosine kinase binding
874	2125	AI102519	c,k		protein

TABLE	A HUMAN	HOWOLOGALE VI	INOTATIONS		th Ally, Docket No. 44921-5039W O
			27/46 28 .877(17:15)		Doc. No. 1798397.1
Seq. ID No. : :=	Identifier	Content Ace./ Ref. Seq. (D No.		Homologous Cene Name	Hemologous Cluster Name
					ESTs, Weakly similar to GEF2_HUMAN GANGLIOSIDE EXPRESSION FACTOR 2 [R.norvegicus], GABA(A) receptor- associated protein-like 2, RIKEN cDNA 0610012F20 gene, ganglioside
875	5967	A1102520	J D W		expression factor 2 ESTs, Weakly similar to GEF2_HUMAN GANGLIOSIDE EXPRESSION FACTOR 2 [R.norvegicus], GABA(A) receptor- associated protein-like 2, RIKEN cDNA 0610012F20 gene, gamma- aminobutyric acid (GABA(A)) receptor- associated protein-like 1, ganglioside expression factor 2
875 876	5969 11563	AI102520 AI102560	p,w General	 	expression factor 2
877	15190	Al102562	b,g,n,p,v		EST, Moderately similar to Cd-7 Metallothionein-2 [H.sapiens], EST, Moderately similar to SMHU1E metallothionein 1E [H.sapiens]
878	19769	AI102570	bb		EST, Highly similar to I49523 Mouse
879	22487	Al102578	General		primary response gene B94 mRNA, 3'end - mouse [M.musculus], Homo sapiens, clone MGC:16332 IMAGE:3842543, mRNA, complete cds, RIKEN cDNA 1200009106 gene, RIKEN cDNA 1600013K19 gene, similar to S. cerevisiae Sec6p and R. norvegicus rsec6, tumor necrosis factor, alpha-induced protein 2
880	19011	AI102618	General	-	
881	23837	AI102620	q,t		
882	23538	Al102727	g,General		solute carrier family 20 (phosphate transporter), member 1, solute carrier family 20 (phosphate transporter), member 2, solute carrier family 20, member 1, solute carrier family 20, member 2
883	17234	Al102741	c	tissue inhibitor of metalloproteinase 3, tissue inhibitor of metalloproteinase 3 (Sorsby fundus dystrophy, pseudoinflammatory)	tissue inhibitor of metalloproteinase 3, tissue inhibitor of metalloproteinase 3 (Sorsby fundus dystrophy, pseudoinflammatory)
884	5891	AI102745	k		
885	6796	Al102753	General		
886	8837	Al102849	o,p		
887	15861	AI102868			phosphoserine aminotransferase
888	3533	AI102877	9		
889	13222	AI102977	General		
890	6806	AI103018	o,u		
891	10659	AI103059	w,cc,General		ATP synthase, H+ transporting, mitochondrial F0 complex, subunit f, iisoform 2, EST, Weakly similar to ATPK_HUMAN ATP SYNTHASE F CHAIN, MITOCHONDRIAL [H.sapiens], ESTs, Highly similar to ATPK_HUMAN ATP SYNTHASE F
892	17400	Al103097	е	<u> </u>	CHAIN, MITOCHONDRIAL [H.sapiens]
893	3584	Al103106	x,aa		
894	13298	AI103143	r		-

TARLES	MANITUMAN!	HOMOLOGUE AN	SKOITATOKI	Market Barrier Co.	Atty : Docket No. 44921-5039W0
UABEES	: InUIVIAIN				Doc. No. 1793397.1
ട്ടത്തി	(M) 17 1	ConBent Ace.	ALC: YES	Homologous Gene	
[y]@	identifier	Ref. Seq. ID No.	Model Code	Name	Homologous Gluster Name
					ESTs, Highly similar to S17516
					hypothetical protein [H.sapiens], ESTs
1					Highly similar to UBIQUITIN- CONJUGATING ENZYME E2-17 KD
					[R.norvegicus], RIKEN cDNA
				i	1110015A16 gene, RIKEN cDNA
1					2610301N02 gene, expressed
					sequence Al327276, ubiquitin
ĺ					conjugating enzyme, ubiquitin-
					conjugating enzyme E2A (RAD6
1					homolog), ubiquitin-conjugating
					enzyme E2B (RAD6 homolog), ubiquitin-conjugating enzyme E2B
					(RAD6 homology), ubiquitin-
					conjugating enzyme E2C, ubiquitin-
895	15981	AI103150	i,x		conjugating enzyme E2G 2
896	3475	Al103245	w		
898	23619	Al103314	р		
899	24181	AI103320	e		
901 902	4355 7622	AI103410 AI103472	General General		
903	20918	AI103472	n		
904	21579	AI103572	General		
905	2222	AI103631	0		
906	2752	AI103641	е		
907	4856	AI103708	ļ <u>.</u>		
908 909	8990 15942	Al103719 Al103738	l,m,y,z		
910	22885	AI103738	e,General		
J		7.1.00025	5,551.6.6.	complement component 4	
					EST, Weakly similar to complement
911	15853	Al103841	x	component 4B	component C4A [H.sapiens]
					EST, Moderately similar to UCRI RAT
					UBIQUINOL-CYTOCHROME C
					REDUCTASE IRON-SULFUR SUBUNIT, MITOCHONDRIAL
1					PRECURSOR [R.norvegicus], EST,
					Weakly similar to UCRI_HUMAN
					UBIQUINOL-CYTOCHROME C
					REDUCTASE IRON-SULFUR
					SUBUNIT, MITOCHONDRIAL
					PRECURSOR [H.sapiens], ESTs,
					Moderately similar to UCRI_HUMAN
					UBIQUINOL-CYTOCHROME C REDUCTASE IRON-SULFUR
					SUBUNIT, MITOCHONDRIAL
					PRECURSOR [H.sapiens], Human
					DNA sequence from clone RP1-228J4
					on chromosome 6 Contains a
					pseudogene similar to UQCRFS1
					(ubiquinol-cytochrome c reductase,
					Rieske iron-sulfur polypeptide 1), ESTs, an STS and GSSs, RIKEN
					cDNA 4430402G14 gene, expressed
					sequence A1875505, ubiquinol-
					cytochrome c reductase, Rieske iron-
912	15050	AI103911	j,y		sulfur polypeptide 1
913	12376	AI103939	u		
914	22271	AI103947	о,у	I	

TABLES	HUMAN	HONOFOGALETY	NOTATIONS		Atty. Docket No. 44921-5039WO Doc. No. 1793397.1
809. [D No. 444	ldentifier	GenBank Ace <i>l</i> Rei. Seq. ID No.	Model Code	Homologous Cene Name	Homologous Gluster Name
915	20833	Al104035	f.q		ESTs, Weakly similar to COXG MOUSE CYTOCHROME C OXIDASE POLYPEPTIDE VIB [M.musculus], Human DNA sequence from clone RP4-591N18 on chromosome 22q13.1-13.2 Contains a COX6B (Cytochrome C Oxidase subunit VIb (EC 1.9.3.1)) pseudogene, ESTs, GSSs and two putative CpG islands, RIKEN cDNA 2010000G05 gene, cytochrome c oxidase subunit VIb
916	7010	AI104099	W		DV57D5640040
917	22101	AI104251	General		DKFZP564O243 protein
918 919 920	22211 10720	Al104258 Al104279 Al104296	General g,m		EST, Weakly similar to IF6_HUMAN EUKARYOTIC TRANSLATION INITIATION FACTOR 6 [H.sapiens], Mus musculus 10 days neonate cerebellum cDNA, RIKEN full-length enriched library, clone:6530402L05, full insert sequence, integrin beta 4 binding protein
921	15416	AI104230	:	<u> </u>	
922	10991	AI104340	a		
923 924	18831 7223	AI104357 AI104373	p e		ESTs, Weakly similar to A29861 actin gamma [H.sapiens], ESTs, Weakly similar to I39393 alpha-actin [H.sapiens], ESTs, Weakly similar to S38782 actin beta' chain [H.sapiens], Homo sapiens mRNA; cDNA DKFZp434B2115 (from clone DKFZp434B2115), RIKEN cDNA 1700052K15 gene, RIKEN cDNA 1700061J02 gene, actin-like 7a, expressed sequence AL023024, expressed sequence AV259599, melanoma X-actin, uncharacterized hypothalamus protein HARP11
925	23574	AI104520	e,g,s	cytochrome c oxidase subunit VIa polypeptide 1, cytochrome c oxidase, subunit VI a, polypeptide 1	
926	18509	AI104528	q		NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 6 (17kD, B17)
927	11680	AI104605	y V		ESTs, Weakly similar to RENAL TRANSCRIPTION FACTOR KID-1 [R.norvegicus], ESTs, Weakly similar to T42682 hypothetical protein DKFZp434G1221.1 [H.sapiens], Homo sapiens, clone MGC:20975 IMAGE:4634585, mRNA, complete cds, expressed sequence AA415813, expressed sequence AI839920, expressed sequence AL024263, hypothetical protein FLJ20531, transcription factor 17, transcription factor 17-like 1, zinc finger protein 91, zinc finger protein homologous to Zfp91 in mouse
929	23689	AI104685	r		
930 931	15377 22957	AI104821 AI104897	o,cc General		hypothetical protein MGC10947, leucine rich repeat (in FLII) interacting protein 2

TABLES	E HUMAN	HOMOLOGUEAN			M. Atty. Docket No. 44921-5089W0 Doc. No. 1793897.1
Seg. ID No.	ldentifer	Cenbank Ace./ Ref. Seq. ID No.		Homologous Cent Name	Homologeus Cluster Name
932	18451	Al104953	o,s		ATP synthase, H+ transporting, mitochondrial F1 complex, delta subunit, EST, Moderately similar to ATPD_HUMAN ATP SYNTHASE DELTA CHAIN, MITOCHONDRIAL PRECURSO [H.sapiens], RIKEN cDNA 0610008F14 gene, expressed sequence AA960090, expressed sequence AI876556, expressed sequence C85518
933	24375	Al104979	n,General		3-oxoacid CoA transferase,
934	18278	AI105080	bb		hypothetical protein FKSG25
935	2196	AI105243	g		
936	5199	Al105272	bb,General		
937 938	12901 7700	AI105301 AI105383	o,s cc,General		
938	13343	AI105383	u		
940	22931	AI105417	e,General		DNA segment, human D4S114, P311 protein
941	23596	AI105435	bb		expressed sequence D17825, glutaryl- Coenzyme A dehydrogenase
942	15893	Al105465	0		ESTs, Highly similar to DHSD_HUMAN SUCCINATE DEHYDROGENASE [H.sapiens], ESTs, Moderately similar to DHSD_HUMAN SUCCINATE DEHYDROGENASE [H.sapiens], succinate dehydrogenase complex, subunit D, integral membrane protein
943	12660	Al111492	C		
944	24211	Al111599 Al111853	General		EST, Moderately similar to 0710252A histone H3 [H.sapiens], ESTs, Highly similar to H33_HUMAN HISTONE H3.3 [H.sapiens], ESTs, Weakly similar to H33_HUMAN HISTONE H3.3 [H.sapiens], ESTs, Weakly similar to JQ1983 H3.3 like histone MH921 - mouse [M.musculus], H3 histone, family 3A, H3 histone, family 3B, H3 histone, family 3B (H3.3B)
946	2539	AI111960	r		
					EGF-containing fibulin-like extracellular matrix protein 1, EGF-containing fibulin like extracellular matrix protein 2, EST, Weakly similar to FBL3_RAT EGF-CONTAINING FIBULIN-LIKE EXTRACELLULAR MATRIX PROTEIN 1 PRECURSOR (FIBULIN-3) (FIBL-3) (T16 PROTEIN) [R.norvegicus], ESTs, Highly similar to FBL3_RAT EGF-CONTAINING FIBULIN-LIKE EXTRACELLULAR MATRIX PROTEIN 1 PRECURSOR (FIBULIN-3) (FIBL-3) (T16 PROTEIN) [R.norvegicus], ESTs, Weakly similar to FBL3_RAT EGF-CONTAINING FIBULIN-LIKE EXTRACELLULAR MATRIX PROTEIN 1 PRECURSOR (FIBULIN-3) (FIBL-3) (T16 PROTEIN) [R.norvegicus], epidermal growth factor-containing fibulin-like extracellular matrix protein
947	5729	AI111990	k		pidermal growth factor-containing fibulin-like extracellular matrix protein 2

TABLE &	MAMUH. S	HOMOPOGALE VI	EXOLIVATIONS		Atty, Docket No. 44921-5089W0 Doc. No. 1793397.1
50g. [D No.	ldentifier	Consont Ace./ Ref. Seq. ID No.	Model Gode	Homologous Cene	Hamologous Cluster Name
948	4049	Al112012	i,q,u,General		EST, Moderately similar to PM17 MOUSE MELANOCYTE PROTEIN PMEL 17 PRECURSOR [M.musculus], Homo sapiens, Similar to glycoprotein (transmembrane) nmb, clone MGC:1696 IMAGE:3345861, mRNA, complete cds, glycoprotein (transmembrane) nmb, silver
949	12908	Al112043	<u> </u>		
950	20041	Al112161	lt		
951	12937	Al112462	General		
952	3713	Al112571	b		
953	12921	Al112636	General		Homo sapiens BAC clone RP11- 335J18 from 2, RIKEN cDNA 1700124F02 gene, expressed sequence Al325217, uridine phosphorylase
954	12965	Al112926	General		
955	7499	Al112986	General		
956	4969	Al113008	r		
957	11817	Al136295	f		DKFZP564O123 protein, putative breast adenocarcinoma marker (32kD)
959	11165	Al136372	С		
960	4045	Al136460	cc		
961	12782	AI136493	k		
962	6850	Al136665	h		ectonucleoside triphosphate diphosphohydrolase 1, ectonucleoside triphosphate diphosphohydrolase 2, ectonucleoside triphosphate diphosphohydrolase 3, ectonucleoside triphosphate diphosphohydrolase 6 (putative function) ESTs, Moderately similar to TISB RAT TIS11B PROTEIN [R.norvegicus], ESTs, Weakly similar to TISB RAT TIS11B PROTEIN [R.norvegicus], butyrate response factor 1, butyrate response factor 1, butyrate response factor 2,
963 964	20920 6552	Al136891 Al137062	p,v	butyrate response factor 1, zinc finger protein, C3H type, 36-like 1	
965	22722	AI137211	i	<u> </u>	
966	13111	AI137224	o,General		hypothetical protein FLJ20260, oxysterol binding protein 2
					DNA segment, Chr 17, ERATO Doi 197, expressed, EST, Weakly similar to ZF37_RAT ZINC FINGER PROTEIN 37 (ZFP-37) [R.norvegicus], ESTs, Weakly similar to I38600 zinc finger protein ZNF135 [H.sapiens], ESTs, Weakly similar to ZINC FINGER PROTEIN ZFP-29 [M.musculus], Homo sapiens GIOT-1 mRNA for gonadotropin inducible transcription repressor-1, partial cds, expressed sequence AI449432, hypothetical protein FLJ14855, zinc finger protein 135 (clone pHZ-17), zinc finger protein 29, zinc finger protein 37, zinc finger
967	15969	AI137302	е		protein homologous to Zfp37 in mouse

TABLE 8	KINAMUH K	HOMOLOGAIS W	SKOTTATIOKE		/Ally. Docket No. 44921-5039
	1 2				Dos. No. 17933
Seq. ID No.	ldentifier	CenBank Ace./ Ref. Seq. ID No.	Westel Goste	Homologous Cene 🐇 Neme	Homologous Gluster Name
NO:	identimen	Iver god in no	ाराकर्वम (ट् कार)	Mame	
969	9166	AI137406	General		protein C receptor, endothelial, pro C receptor, endothelial (EPCR)
					EST, Weakly similar to ZF37_RAT
970	9525	Al137516			ZINC FINGER PROTEIN 37 (ZFP-[R.norvegicus], ESTs, Weakly simit b B32891 finger protein 2, placent [H.sapiens], ESTs, Weakly similar MLZ4 MOUSE ZINC FINGER PROTEIN MLZ-4 [M.musculus], Hc sapiens cDNA FLJ14967 fis, clone THYRO1000242, moderately similar ZINC FINGER PROTEIN 84, expressed sequence Al854635, zin finger protein 113, zinc finger protein 268, zinc finger protein 37, zinc finger protein 46, zinc finger protein 84 (HPF2), zinc finger protein homologous to Zfp37 in mouse
971	6638	Al137579	General		nonologous to zipsi in mouse
					EST, Weakly similar to IMB3_HUM IMPORTIN BETA-3 SUBUNIT [H.sapiens], Homo sapiens cDNA FLJ12978 fis, clone NT2RP200632 RAN binding protein 6, karyopherin (importin) bata 3
972	7414	Al137586	General		(importin) beta 3 EST, Highly similar to R5HU7 ribosomal protein L7, cytosolic
					[H.sapiens], EST, Weakly similar to RL7 MOUSE 60S RIBOSOMAL PROTEIN L7 [M.musculus], ESTs, Highly similar to R5HU7 ribosomal protein L7, cytosolic [H.sapiens], calponin like transmembrane doma
973	11321	Al137752	z		protein, ribosomal protein L7
974	23473	AI137932	!		
975	13158	AI138024	1		UDP-glucose ceramide
976	13467	AI138034	cc		glucosyltransferase
977	11377	Al138105	v		g.accoyia.anorciacc
978	6790	Al144801	d,h		
979	6506	Al144919	j.l,y		
980	8027	Al144958	i		
982	14458	AI145095	General		
983 984	7476 17545	AI145202 AI145384	g		
984 985	17479	AI145384 AI145385	e r		
303	4194	AI145387	<u></u>		
986		AI145722	g		
	18634	M1143/22			
	8634 8339	AI145722 AI145761	y,General		
987 988					RIKEN cDNA 2610020J05 gene, pseudouridine synthase 1, pseudouridylate synthase 1
989	2059	Al145761 Al146005	y,General h,General		pseudouridine synthase 1, pseudouridylate synthase 1 translocase of inner mitochondrial membrane 10 homolog (yeast), translocase of inner mitochondrial
987 988 989 990	2059 23224 5232	Al145761 Al146005 Al146033 Al168942	y,General h,General o	branched chain keto acid dehydrogenase E1, beta polypeptide (maple syrup urine disease), branched chain ketoacid dehydrogenase E1, beta polypeptide	pseudouridine synthase 1, pseudouridylate synthase 1 translocase of inner mitochondrial membrane 10 homolog (yeast),
987 988 989 990 991 992	2059 23224 5232 18472	Al145761 Al146005 Al146033 Al168942 Al168975	y,General h,General o	dehydrogenase E1, beta polypeptide (maple syrup urine disease), branched chain ketoacid dehydrogenase E1, beta	pseudouridine synthase 1, pseudouridylate synthase 1 translocase of inner mitochondrial membrane 10 homolog (yeast), translocase of inner mitochondrial
987 988 989 990	2059 23224 5232	Al145761 Al146005 Al146033 Al168942	y,General h,General o	dehydrogenase E1, beta polypeptide (maple syrup urine disease), branched chain ketoacid dehydrogenase E1, beta	pseudouridine synthase 1, pseudouridylate synthase 1 translocase of inner mitochondrial membrane 10 homolog (yeast), translocase of inner mitochondrial

TABLES	E HUMAN	HOMOLOGUEAN	EXIONATION		Atty. Docket No. 44921-5089WO Doc. No. 1793897.1
Seq. ID No.	ldentiffer	Genisank Ace./ Ref. Seq. (D No.	Model Gode	Hemelogous Ceme	Homologous Queter Name
995	17386	Al169144	o		
996	10984	Al169156	o,u		
997	8205	Al169176	е		
998	12979	Al169177	е		immediate early response 3
000	2607	A1460244			Homo sapiens clone 24468 mRNA sequence, Mus musculus 0 day neonate skin cDNA, RIKEN full-length enriched library, clone:4633401122, full insert sequence, heterogeneous nuclear ribonucleoprotein C, heterogeneous nuclear ribonucleoprotein C (C1/C2), hnRNP-associated with lethel valley.
999	2607	AI169211	С		associated with lethal yellow ATPase, H+ transporting, lysosomal (vacuolar proton pump), subunit 1, EST, Weakly similar to I54197 hypothetical protein [H.sapiens], ESTs, Weakly similar to VAS1_RAT VACUOLAR ATP SYNTHASE SUBUNIT S1 PRECURSOR (V- ATPASE S1 SUBUNIT) (V-ATPASE S1 ACCESSORY PROTEIN) (V- ATPASE AC45 SUBUNIT) (C7-1 PROTEIN) [R.norvegicus], Homo sapiens cDNA FLJ12563 fis, clone NT2RM4000820, weakly similar to VACUOLAR ATP SYNTHASE SUBUNIT AC45 PRECURSOR (EC
1000	22661	Al169265	s,z		3.6.1.34)
1001	13239	Al169278	g,j,l,y,z	,	0.011.017
1002	24162	Al169279	m		
1003	16879	AI169284	0		ADP-ribosylation factor-like 6 interacting protein
1004	24213	Ai169289	p		EST, Moderately similar to 0710252A histone H3 [H.sapiens], ESTs, Highly similar to H33_HUMAN HISTONE H3.3 [H.sapiens], ESTs, Weakly similar to H33_HUMAN HISTONE H3.3 [H.sapiens], ESTs, Weakly similar to JQ1983 H3.3 like histone MH921 - mouse [M.musculus], H3 histone, family 3A, H3 histone, family 3B, H3 histone, family 3B (H3.3B)
1005	13240	Al169311	cc		
1006	5931	AI169324	b		ļ
1007	20891	AI169337	d		hypothetical protein
1008	11979	Al169365 Al169372	s		EST, Weakly similar to S13101 cytochrome P450 c117 - rat [R.norvegicus], RIKEN cDNA 2010301M18 gene, RIKEN cDNA 2010318C06 gene, RIKEN cDNA 2210009K14 gene, cytochrome P450, 2c29, cytochrome P450, subfamily IIC (mephenytoin 4-hydroxylase), polypeptide 19

	e e casar	HOMOLOGUEAN	SKOTATOKK		Atty, Docket No. 44921-5089W0 Doc. No. 1798397.1
Seg. ID No.	ldentifier	ConEank Acc./ Ref. Seq. ID No.	Model Gode	Homologous Ceine Name	Homologous Cluster Name
1010	20697	Al169494	o,u		ATPase, H+ transporting, lysosomal (vacuolar proton pump), 42 kDa, ATPase, H+ transporting, lysosomal (vacuolar proton pump), member D, ESTs, Moderately similar to VAOD_HUMAN VACUOLAR ATP SYNTHASE SUBUNIT D (V-ATPASE D SUBUNIT) (VACUOLAR PROTON PUMP D SUBUNIT) (V-ATPASE AC39 SUBUNIT) (V-ATPASE 40 KDA ACCESSORY PROTEIN) (P39) [H.sapiens]
1011	8234	AI169517	Z		
1012	18343	AI169648	0		
1013	10839	AI169655	I,m		
1014	24146	AI169668	j ,l		ATP-binding cassette, sub-family F (GCN20), member 1, ATP-binding cassette, sub-family F (GCN20), member 2, hypothetical protein FLJ11198
1015	22575	Al169728	r		EST, Highly similar to T47184 hypothetical protein DKFZp434F1526.1 [H.sapiens], ESTs, Weakly similar to T47184 hypothetical protein DKFZp434F1526.1 [H.sapiens], hypothetical protein FLJ10889
1016	804	AI169756	cc		Gene 33/Mig-6, RIKEN cDNA 1300002F13 gene
1017	8213	Al169883	р	ferritin light chain 1, ferritin, light polypeptide	ESTs, Highly similar to FRHUL ferritin light chain [H.sapiens], ESTs, Moderately similar to FRHUL ferritin light chain [H.sapiens], PR00470 protein, RIKEN cDNA 4933416E14 gene, ferritin light chain 2, ferritin, light polypeptide
1018	3916	AI169947	i,bb		
1019	3733	AI170053	u,General		
1020	14179	AI170224	cc		
1022	3547	Al170263 Al170279	r General		interleukin 20 receptor, alpha ESTS, Weakly similar to ZNT4_HUMAN ZINC TRANSPORTER 4 [H.sapiens], RIKEN cDNA 1810059J10 gene, hypothetical protein DKFZp547M236, hypothetical protein FLJ12496, solute carrier family 30 (zinc transporter), member 1, solute carrier family 30 (zinc transporter), member 4
					PDZ and LIM domain 1 (elfin), PDZ and LIM domain 3, PDZ-LIM protein mystique, RIKEN cDNA 1110003B01
4022	11524	A1470240			gene, Rattus norvegicus LIM-domain protein LMP-1 mRNA, complete cds, Z-band alternatively spliced PDZ-motif, actinin alpha 2 associated LIM protein, alpha-actinin-2-associated LIM protein,
1023	11524	Al170340	j,y,z		protein LMP-1 mRNA, complete cds, Z- band alternatively spliced PDZ-motif, actinin alpha 2 associated LIM protein,
1024	2729	Al170363	j,y,z e,i		protein LMP-1 mRNA, complete cds, Z- band alternatively spliced PDZ-motif, actinin alpha 2 associated LIM protein, alpha-actinin-2-associated LIM protein,
					protein LMP-1 mRNA, complete cds, Z- band alternatively spliced PDZ-motif, actinin alpha 2 associated LIM protein, alpha-actinin-2-associated LIM protein,

			HOMOLOGUEAN			Atty. Docket No. 44921-5039W0 Doc. No. 1798397.1
Bike 2, RIKEN CDNA 610012F26 1028 5958	Seq. ID No. +	ldentifier	Geneenk Aced Ref. Seq. ID No.	Model Code	Homologous Cans Nemo	Homologous Gluster Name
1039 9757 Al170693 b	1028	5968	A1170692	y,aa		I
1031		9757		b		
1031	1030	18905	Al170770	e,s		
Wolffian duct includes surrounding region cDNA (ERM full-length enriched library, clone:6720486F14, https://doi.org/10.1002/10.24 gene, TRAF associated motility receptor (RHAMM), hyaluronan-mediated motility receptor (RHAMM), hyaluronan mediated motility receptor (RHAMM), hyaluronan mediated motility receptor (RHAMM), hyaluronan mediated motility receptor (RHAMM), hyaluronan-mediated motility rec	1031	16170	AI170894	i		RIKEN cDNA 2310076L09 gene, adipose differentiation related protein, adipose differentiation-related protein
1034 13285					motility receptor (RHAMM), hyaluronan-mediated	wolffian duct includes surrounding region cDNA, RIKEN full-length enriched library, clone:6720466F14, full insert sequence, RIKEN cDNA 0610027D24 gene, TRAF4 associated factor 1, hyaluronan mediated motility receptor (RHAMM), hyaluronan-
1034 13285	1033	17391	A117 1354	D		heterogeneous nuclear
EST, Moderately similar to NUAM_HUMAN NADH-UBIQUINONE NUAM_HUMAN NADH-UBIQUINONE NADH-UBIQUINO	1034	13285	A1171361	h		
RIKEN cDNA 2010107E04 gene, chromosome 14 open reading frame expressed sequence AU043134, expressed sequence AV124504						NUAM_HUMAN NADH-UBIQUINONE OXIDOREDUCTASE 75 KD SUBUNIT PRECURSOR [H.sapiens], NADH dehydrogenase (ubiquinone) Fe-S protein 1 (75kD) (NADH-coenzyme Q
Chromosome 14 open reading frame expressed sequence AU043134, expressed sequence AV124504	1036	18126	AI1/1369	W		DU(51 - D11 - 0040407504
EST, Moderately similar to N12M_HUMAN NADH-UBIQUINONE OXIDOREDUCTASE B22 SUBUNIT [H.sapiens], NADH dehydrogenase (Ubiquinone) 1 beta subcomplex, 9 (22kD, B22)						chromosome 14 open reading frame 2 expressed sequence AU043134,
1041 21183						NI2M_HUMAN NADH-UBIQUINONE OXIDOREDUCTASE B22 SUBUNIT [H.sapiens], NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 9
ESTs, Highly similar to FRHUL ferritin light chain [H.sapiens], ESTs, Moderately similar to FRHUL ferritin light chain [H.sapiens], PRO0470 protein, RIKEN cDNA 2010009K05 gene, RIKEN cDNA 4933416E14 gene, cysteine conjugate-beta lyase; cytoplasmic (glutamine transaminase K, kyneurenine aminotransferase), ferritin light chain 1, ferritin, ferritin light chain 2, ferritin, light polypeptide, hypothetical protein 669 1043 11437 Al171794 i polypeptide polypeptide, hypothetical protein 669 1044 2625 Al171800 cc 1045 23579 Al171802 v 1046 11708 Al171807 I,t ATP synthase, H+ transporting, mitochondrial F1 complex, epsilon subunit, RIKEN cDNA 2410043G19 gene, expressed sequence AV00064	1040	15345	AI171587	1		
1042 8215 Al171692 i light polypeptide polypeptide, hypothetical protein 669 1043 11437 Al171794 i Polypeptide polypeptide, hypothetical protein 669 1044 2625 Al171800 cc	1041	21183	Al171676	k		Moderately similar to FRHUL ferritin light chain [H.sapiens], PRO0470 protein, RIKEN cDNA 2010009K05 gene, RIKEN cDNA 4933416E14 gene, cysteine conjugate-beta lyase; cytoplasmic (glutamine transaminase
1043 11437 Al171794 i 1044 2625 Al171800 cc 1045 23579 Al171802 v 1046 11708 Al171807 I,t ATP synthase, H+ transporting, mitochondrial F1 complex, epsilon subunit, RIKEN cDNA 2410043G19 1047 17204 Al171844 s,y,z gene, expressed sequence AV00064						ferritin light chain 2, ferritin, light
1044 2625 AI171800 cc 1045 23579 AI171802 v 1046 11708 AI171807 I,t ATP synthase, H+ transporting, mitochondrial F1 complex, epsilon subunit, RIKEN cDNA 2410043G19 1047 17204 AI171844 s,y,z gene, expressed sequence AV00064				i	light polypeptide	polypeptide, hypothetical protein 669
1045 23579 Al171802 v 1046 11708 Al171807 I,t ATP synthase, H+ transporting, mitochondrial F1 complex, epsilon subunit, RIKEN cDNA 2410043G19 1047 17204 Al171844 s,y,z gene, expressed sequence AV00064				li .		
1046 11708 Al171807 I,t ATP synthase, H+ transporting, mitochondrial F1 complex, epsilon subunit, RIKEN cDNA 2410043G19 1047 17204 Al171844 s,y,z gene, expressed sequence AV00064						
ATP synthase, H+ transporting, mitochondrial F1 complex, epsilon subunit, RIKEN cDNA 2410043G19 1047 17204 AI171844 s,y,z gene, expressed sequence AV00064					<u> </u>	
						mitochondrial F1 complex, epsilon subunit, RIKEN cDNA 2410043G19
	1047 1048	17204 4420	AI171844 AI171916	s,y,z m	 	gerie, expressea sequence AV000645

TABLES	BE CHŪMAN	HOMOLOGUE AN	EMOTATIONS		Atty. Docket No. 44921-5039WO Doc. No. 1793897.1
809. [D No	ldenliiter	Cenbank Acc./ Ref. Seq. ID No.	Model Gode	Homologous Cene. Name	Homologous Cluster Namo
1049 1050	3266 19012	Al171948 Al172056	I,m	international participation and	DKFZP564F0522 protein, ESTs, Weakly similar to T08675 hypothetical protein DKFZp564F0522.1 [H.sapiens]
1050	11205	AI172057	a,q,bb		
1052	6057	Al172102	b		_
1053	19128	Al172103	m		
1054	15673	Al172107	z		KIAA1883 protein, sirtuin (silent mating type information regulation 2, S.cerevisiae, homolog) 2, sirtuin 2 (silent mating type information regulation 2, homolog) 2 (S.cerevisiae)
1055	6630	Al172184	n		Alpha-fetoprotein, ESTs, Weakly
1056	11968	AI172208	bb		similar to ALPHA-FETOPROTEIN PRECURSOR [M.musculus], ESTs, Weakly similar to FPHU alpha- fetoprotein precursor [H.sapiens], alpha fetoprotein, alpha-fetoprotein
1057	6974	AI172263	l,m		
1058	23313	Al172271 Al172272	d General		ESTs, Highly similar to JC4577 transcription elongation factor T1 [H.sapiens], expressed sequence Al326274, transcription elongation factor A (SII), 2
1060	15382	AI172302	I,p,General		lactor A (on), 2
1061	18689	Al172329	l		
1062	17887	Al172414	0		
1063	3042	Al172447	General		ESTs, Highly similar to BCL3 [M.musculus], ESTs, Weakly similar to I-kappa B alpha chain [M.musculus], hypothetical protein MGC15396, nuclear factor of kappa light chain gene enhancer in B-cells inhibitor, alpha, nuclear factor of kappa light chain gene enhancer in B-cells inhibitor, beta, nuclear factor of kappa light polypeptide gene enhancer in B-cells inhibitor, alpha, nuclear factor of kappa light polypeptide gene enhancer in B-cells inhibitor, epsilon, testisspecific ankyrin motif containing protein
1003	10042	74172447	Contrai		ESTs, Highly similar to ISOCITRATE
1064	17291	Al172491	bb		DEHYDROGENASE [R.norvegicus], Isocitrate dehydrogenase 1, soluble, isocitrate dehydrogenase 1 (NADP+), soluble, isocitrate dehydrogenase 2 (NADP+), mitochondrial
1065	26222	Al172506	р		
1066	13095	AI172595	r		
1067	8795 6454	Al172618 Al175342	General j,l,m,y		BACULOVIRAL IAP REPEAT- CONTAINING PROTEIN 6 (UBIQUITIN-CONJUGATING BIR- DOMAIN ENZYME APOLLON) [H.sapiens], ESTs, Moderately similar to T31067 BIR repeat containing ubiquitin-conjugating enzyme BRUCE - mouse [M.musculus], baculoviral IAP repeat-containing 6, hypothetical protein FLJ13855, likely ortholog of mouse ubiquitin-conjugating enzyme E2-230K

	lentifier	GenBank Acc./ Rof. Seq. ID No.	Model Gode	Homologous Cene Namo	Atty, Docket No. 44921-5089We Doc, No. 1798397.1 Homologous Gluster Name EST, Highly similar to RASN RAT TRANSFORMING PROTEIN P21/N- RAS [R.norvegicus], EST, Weakly similar to TVHURR transforming protein R-ras [H.sapiens], Harvey rat sarcoma oncogene, subgroup R, RIKEN cDNA 2610016H24 gene, RIKEN cDNA 4930526B11 gene, Ris, expressed sequence Al573426, neuroblastoma RAS viral (v-ras) oncogene homolog, neuroblastoma ras oncogene, related RAS viral (r-ras) oncogene homolog, similar to mouse
1070 44	445	Al175466	x		TRANSFORMING PROTEIN P21/N-RAS [R.norvegicus], EST, Weakly similar to TVHURR transforming protein R-ras [H.sapiens], Harvey rat sarcoma oncogene, subgroup R, RIKEN cDNA 2610016H24 gene, RIKEN cDNA 4930526B11 gene, Ris, expressed sequence Al573426, neuroblastoma RAS viral (v-ras) oncogene homolog, neuroblastoma ras oncogene, related RAS viral (r-ras)
1070 44	445	Al175466	×		RIKEN cDNA 4930526B11 gene, Ris, expressed sequence Al573426, neuroblastoma RAS viral (v-ras) oncogene homolog, neuroblastoma ras oncogene, related RAS viral (r-ras)
1070 44	445	A1175466	x		
				1	Ras, dexamethasone-induced 1
l l					ESTs, Weakly similar to NHPX RAT NHP2/RS6 FAMILY PROTEIN YEL026W HOMOLOG [R.norvegicus], RIKEN cDNA 2410130M07 gene, non-histone chromosome protein 2 (S. cerevisiae)-like 1, nucleolar protein family A, member 2 (H/ACA small
1071 34	418	Al175475	m		nucleolar RNPs), sperm specific antigen 1 ESTs, Highly similar to S25432
1072 18	8507	AI175551	bb		translation elongation factor eEF-1 beta chain [H.sapiens], eukaryotic translation elongation factor 1 beta 2
	0217	Al175628	w		translation elongation ractor i beta 2
	262	AI175833	i,m,x		
		AI175875	r		
		Al175959	I,General		
1077 70	022	AI176041	h,n		EST, Moderately similar to TIG2_HUMAN RETINOIC ACID RECEPTOR RESPONDER PROTEIN 2 PRECURSOR [H.sapiens], retinoic acid receptor responder (tazarotene
1078 21	1467	AI176061	t		induced) 2
	8581	AI176160	General		
	4159	AI176169	g		
1081 21	1742	Al176172	w		
1082 10	0182	Al176185	v	FBJ osteosarcoma oncogene, v-fos FBJ murine osteosarcoma viral oncogene homolog	
		Al176265	General		
					EST, Moderately similar to T02747 phospholipid-hydroperoxide glutathione peroxidase [H.sapiens], EST, Weakly similar to T02747 phospholipid-hydroperoxide glutathione peroxidase [H.sapiens], ESTs, Weakly similar to GSHH RAT PHOSPHOLIPID HYDROPEROXIDE GLUTATHIONE PEROXIDASE [R.norvegicus], Homo sapiens
1084 69	905	Al176275	a		PRO2893 mRNA, complete cds, RIKEN cDNA 2310016C16 gene, RIKEN cDNA 3110050F08 gene, glutathione peroxidase 4, glutathione peroxidase 4 (phospholipid hydroperoxidase)
1004 09	J-J-J-	7.170210	<u> </u>		Homo sapiens cDNA FLJ12570 fis, clone NT2RM4000895, UDP-N-acteylglucosamine pyrophosphorylase
1085 12	2999	AI176276	cc		1

		HOMOLOGUEAA		La Com Spirit	Atty. Docket No. 4/1921-5039W0 Doc. No. 1793397.1
		GenBank Ace./ Ref. Seq. ID No.		Homologous Cene Name	Homologous Gluster Namo
					ESTs, Weakly similar to B Chain B, Crystal Structure Of The D1d2 Sub- Complex From The Human Snrnp Core Domain [H.sapiens], small nuclear ribonucleoprotein D2
1086	16438	Al176294	е		polypeptide (16.5kD)
1087	21130	AI176298	у		
1088 1089	3014	AI176362 AI176363	e r		
1090	15015 19006	AI176393	x		
1091	20001	Al176396	0		ESTs, Highly similar to C560_HUMAN SUCCINATE DEHYDROGENASE CYTOCHROME B560 SUBUNIT, MITOCHONDRIAL PRECURSOR [H.sapiens], succinate dehydrogenase complex, subunit C, integral membrane protein, 15kD
1092	12174	Al176435	j,m		
1093	15191	Al176456	b,o,t,v,cc		ESTs, Moderately similar to AF078844 1 hqp0376 protein [H.sapiens], expressed sequence AA409533
1094	24236	AI176473	d,General		ESTs, Highly similar to HS9B RAT HEAT SHOCK PROTEIN HSP 90-BETA [R.norvegicus], Mus musculus, clone IMAGE:3584589, mRNA, partial cds, expressed sequence C81438, heat shock 90kD protein 1, beta, heat shock protein, 84 kDa 1, heat shock
1095	16518	AI176546	V		protein, 86 kDa 1
1096	12436	AI176592	General		ESTs, Weakly similar to SYC_HUMAN CYSTEINYL-TRNA SYNTHETASE [H.sapiens], cysteinyl-tRNA synthetase, hypothetical protein FLJ12118
1097	2536	AI176616	I,v,General		FLJ12110
1099	18525	Al176792	u		
1100	23449	AI176828	g		
1101	23299	AI176839	General	<i>"</i>	
1102	3580	Al176848	е		
1103	22103	AI176849	d,General		
1104	16036	AI176855	f .		
1105 1106	15588 16917	Al176916 Al176951	General t		phosphomannomutase 1
1107	16124	Al176963	cc		with Glu/Asp-rich carboxy-terminal domain, 2, Cbp/p300-interacting transactivator, with Glu/Asp-rich carboxy-terminal domain, 4, ESTs, Weakly similar to MRG1_HUMAN MSG-RELATED PROTEIN 1 [H.sapiens], expressed sequence AW742964
1108	15146	AI176969	b,General		
1109	5786	AI177058	1		
1110 1112	2852 3156	AI177059 AI177092	g g		
1113	14384	Al177096	a		EST, Moderately similar to APT RAT ADENINE PHOSPHORIBOSYLTRANSFERASE [R.norvegicus], Mouse adenine phosphoribosyltransferase (APRT), complete cds, adenine phosphoribosyltransferase, adenine phosphoribosyltransferase, expressed sequence C85684

TABLES	HUMAN	HOMOLOGUE AN	NOTATIONS		Atty, Docket No. 44921-5089Wo
					Doc. No. 1798397.1
Scq. ID No.	ldenWifer	GenBank Acc./ Ref. Seq. (D No.	Model Code	Homologous Cane	Homologous Gluster Name
100					ESTs, Weakly similar to COMPLEMENT C1Q
					SUBCOMPONENT, C CHAIN
					PRECURSOR [M.musculus], Homo
					sapiens, Similar to complement
					component 1, q subcomponent, c polypeptide, clone MGC:17279
					IMAGE:4212772, mRNA, complete
					cds, complement component 1, q subcomponent, beta polypeptide,
					complement component 1, q
1114	13310	AI177119	General		subcomponent, c polypeptide, expressed sequence Al385742
1115	24049	AI177341	g,p,s,u		CGI-10 protein
1116	15964	AI177360	o,General		
					ESTs, Highly similar to B27079 fibronectin receptor beta chain
					precursor [H.sapiens], integrin beta 1
					(fibronectin receptor beta), integrin
					beta 2, integrin, beta 1 (fibronectin
					receptor, beta polypeptide, antigen CD29 includes MDF2, MSK12),
					integrin, beta 2 (antigen CD18 (p95),
					lymphocyte function-associated
1117	14989	A1477266			antigen 1; macrophage antigen 1 (mac-
1118	7975	AI177366 AI177374	u aa		1) beta subunit)
					dickkopf (Xenopus laevis) homolog 3,
	0000	A1477005			dickkopf (Xenopus laevis) homolog 4,
1119	3006	AI177395	k		soggy-1 gene ESTs, Highly similar to ROA3 HUMAN
					HETEROGENEOUS NUCLEAR
					RIBONUCLEOPROTEIN A3
					[H.sapiens], ESTs, Highly similar to S12520 core protein A1 [H.sapiens],
					ESTs, Weakly similar to ROA2
					MOUSE HETEROGENEOUS
					NUCLEAR RIBONUCLEOPROTEINS
					A2/B1 [M.musculus], Human DNA sequence from clone 522P13 on
					chromosome 6p21.31-22.3. Contains a
]					60S Ribosomal Protein L21
					pseudogene and an HNRNP A3
					(Heterogenous Nuclear Riboprotein A3, FBRNP) pseudogene. Contains
					ESTs, STSs and GSSs, RIKEN cDNA
					3010025E17 gene, Ras-GTPase-
					activating protein SH3-domain binding protein, heterogeneous nuclear
					ribonucleoprotein A1, heterogeneous
					nuclear ribonucleoprotein A2/B1,
1120 1121	17570 9521	Al177683 Al177706	r b		hypothetical protein 23851
1141	3321	<u></u>	<u> </u>	<u> </u>	ESTs, Moderately similar to
					PBEF_HUMAN PRE-B CELL
					ENHANCING FACTOR PRECURSO
1122	14425	Al177755	g.General		[H.sapiens], pre-B-cell colony- enhancing factor
1123	10611	Al177790	j,m		oa.ong radio
1124	5356	Al177813	cc		modulator recognition factor I
1125	11791	AI177843	General		sarcoma amplified sequence

TABLE 8	RIMMAN)	HOWOTOGME W	NOTATIONS I		Any, Docket No. 44921-5039WO
					Doc. No. 1793997.1
Seq. (D No.	Odemiiiter	GonBook Ace./ Ref. Seq. ID No.	Model Gode	Homologous Cene Neme	Homologous Civeter Name
					EH-domain containing 3, EH-domain containing 4, ESTs, Highly similar to EP15 MOUSE EPIDERMAL GROWTH FACTOR RECEPTOR SUBSTRATE SUBSTRATE 15 [M.musculus], Homo sapiens CDNA FLJ13682 fis, clone
					PLACE2000015, weakly similar to EPIDERMAL GROWTH FACTOR RECEPTOR SUBSTRATE SUBSTRATE 15, Mus musculus adult male cecum cDNA, RIKEN full-length enriched library, clone:9130014G17, full insert sequence, epidermal growth factor receptor pathway substrate 15, epidermal growth factor receptor
1126	14484	AI177867	General		substrate EPS15R
1127	5780	Al177869	General		EST, Weakly similar to TESTIN 2 [M.musculus], LIM and cysteine-rich domains 1, four and a half LIM domains 2, testis derived transcript, testis derived transcript (3 LIM domains), vascular Rab-GAP/TBC- containing
1121	3700	A1177005	General		Containing
					ESTs, Weakly similar to TGIF MOUSE 5'-TG-3' INTERACTING FACTOR [M.musculus], Homo sapiens TGF beta induced transcription factor 2-like mRNA, partial sequence, RIKEN cDNA 5430405H02 gene, RIKEN cDNA 5730599009 gene, TG interacting factor, TGFB-induced factor (TALE
1128	19184	AI178025	General		family homeobox), TGFB-induced factor 2 (TALE family homeobox)
					ESTs, Highly similar to T13963 formin related protein, lymphocyte specific - mouse [M.musculus], ESTs, Moderately similar to T13963 formin related protein, lymphocyte specific - mouse [M.musculus], Homo sapiens mRNA; cDNA DKFZp762B245 (from clone DKFZp762B245); partial cds,
1129	6059	AI178245	c,General		KIAA1902 protein, formin-like
1130 1131	23248 4073	AI178267 AI178272	у 0		chromosome 1 open reading frame 9
1131	7838	AI178291	e		
1133	18996	AI178326	у		
					EST, Highly similar to I49523 Mouse primary response gene B94 mRNA, 3'end - mouse [M.musculus], Homo sapiens, clone MGC:16332 IMAGE:3842543, mRNA, complete cds, RIKEN cDNA 1200009106 gene, RIKEN cDNA 1600013K19 gene, similar to S. cerevisiae Sec6p and R. norvegicus rsec6, tumor necrosis
1134	22488	AI178392	b		factor, alpha-induced protein 2
1135 1136	18800 22197	AI178504 AI178527	n,p,aa g,General		-
1100	12101	7.117.0021	3,30110101		ESTs, Weakly similar to MCM3_HUMAN DNA REPLICATION LICENSING FACTOR MCM3 [H.sapiens], minichromosome
1137	3401	AI178684	bb		maintenance deficient (S. cerevisiae) 3
1138	17713	AI178700	m		
1139	14874	AI178735	e	L	

TABLES	E HUMAN	HOMOROGAE W	EXOTATIONS		Aii y, Docket No. 44921-5939W0 Doc, No. 1793397.1
Seg, ID No.	lden/lifier	GenBenk Acc./ Ref. Seq. ID No.	Model Gode	Homologous Gene Name	Homologous Cluster Name
1140	23567	AI178746	v,General	The state of the s	
1141	18907	Al178971	С		EST, Moderately similar to HART1 hemoglobin alpha-1 chain - rat [R.norvegicus], EST, Weakly similar to A45964 hemoglobin alpha chain - mouse [M.musculus], ESTs, Moderately similar to HART1 hemoglobin alpha-1 chain - rat [R.norvegicus], Hemoglobin, alpha 1, hemoglobin alpha, adult chain 1, hemoglobin, alpha 1, hemoglobin, alpha 1, hemoglobin, alpha 1, hemoglobin, alpha 2
1142	20991	Al178979	i		
1143	5887	Al179099	q,t		biotinidase, vanin 1, vanin 2, vanin 3
1144	8477	AI179167	b,e,General		
1145	3348	AI179288	u,v		
1146 1147	13608 8849	AI179314 AI179315	g,p		
1148	13611	Al179378	v,General m,x	collagen, type V, alpha 2, procollagen, type V, alpha 2	EST, Weakly similar to MAST CELL PROTEASE 7 PRECURSOR [M.musculus], ESTs, Weakly similar to MCT7 RAT MAST CELL PROTEASE 7 PRECURSOR [R.norvegicus], RIKEN cDNA 2410039E18 gene, RIKEN cDNA 4933401F05 gene, marapsin, mast cell protease 7, protease, serine, 21 (testisin), protease, serine, 22, protease, serine, 8 (prostasin), tryptase delta 1, tryptase, alpha EST, Highly similar to CA13_HUMAN COLLAGEN ALPHA 1(III) CHAIN PRECURSOR [H.sapiens], EST, Weakly similar to CA25_HUMAN COLLAGEN ALPHA 2(V) CHAIN PRECURSO [H.sapiens], EST, Weakly similar to I49607 procollagen type V alpha 2 - mouse [M.musculus], RIKEN cDNA 2810002D19 gene, collagen, type III, alpha 1 (Ehlers-Danlos syndrome type IV, autosomal dominant), collagen, type V, alpha 2, hypothetical protein DKFZp434F0318, hypothetical protein MGC12921, procollagen, type VIII, alpha 1, procollagen, type V, alpha 2, procollagen, type V, alpha 2, procollagen, type V, alpha 2, procollagen, type V, alpha 2, procollagen, type V, alpha 2, procollagen, type V, alpha 2, procollagen, type VIII, alpha 1
1150	13614	AI179407	e,t,General		procedugeri, type xiii, espire :
1151	15042	AI179422	b,General		
1152	2768	Al179481	i,General		
1153	24041	AI179580	b,i		
1154	19822	Al179599	o,General		
1155	23270	Al179601	q,General		
1156	5901	Al179605	е		
1157 1158	16081 14564	Al179610 Al179717	g,i,p k	heme oxygenase (decycling) 1	heme oxygenase (decycling) 1
1159	7918	AI179750	General		
1160	9097	Al179795 Al179875	g o,General	hypothetical protein similar to mouse aldehyde reductase 6 (renal), renal- specific oxido-reducatse	

TABLE	B: HUMAN	HONOLOGAE VI	BKÖNTÄTTÖKN		Atty: Docket No. 4/1921-5039W0 Doc. No. 1793397.1
Seg. (D No.	ldentifier	Consenk Acel Ref. Seq. 10 No.	Model Code	Homologous Gene* Name	Homologous Cluster Name
1162	23989	Al179953	a		gap junction membrane channel protein beta 2, gap junction membrane channel protein beta 6, gap junction protein, beta 2, 26kD (connexin 26), gap junction protein, beta 6 (connexin 30)
1163	12899	AI179967	b		
1164	1687	Al179971	c	hemoglobin alpha, adult chain 2, hemoglobin, alpha 1	EST, Moderately similar to HART1 hemoglobin alpha-1 chain - rat [R.norvegicus], ESTs, Moderately similar to HART1 hemoglobin alpha-1 chain - rat [R.norvegicus], RIKEN cDNA 2510042H12 gene, hemoglobin alpha, adult chain 1, hemoglobin, alpha 1, hemoglobin, alpha 2
1165	22569	Al179979	General		<u> </u>
1166	23514	AI179986	o,General		phosphoserine phosphatase, phosphoserine phosphatase-like
1167	15892	AI179988	c,General		
1168	12402	AI180004	g		DnaJ (Hsp40) homolog, subfamily B,
1169	5443	AI180165	General		member 3, DnaJ (Hsp40) homolog, subfamily B, member 6, ESTs, Moderately similar to HSJ2_HUMAN DNAJ PROTEIN HOMOLOG 2 [H.sapiens], ESTs, Weakly similar to HSJ2_HUMAN DNAJ PROTEIN HOMOLOG 2 [H.sapiens], Homo sapiens, clone IMAGE:3930327, mRNA, partial cds, RIKEN cDNA 2810451A06 gene, expressed sequence AU020082
1170	5481	AI180170	General		eukaryotic translation termination factor 1
1171	24028	AI180239	1		
1172	17089	AI180281	g		
1173	3701	AI180306			KIAA0273 gene product, KIAA1796 protein
1174	3352	AI180334	m aa		protein
1175	24368	Al180392	I,m		ESTs, Highly similar to NBP_HUMAN NUCLEOTIDE-BINDING PROTEIN [H.sapiens], hypothetical protein FLJ12660, nucleotide binding protein 1 (E.coli MinD like), nucleotide binding protein 2 (E.coli MinD like)
1176	14337	Al180414 Al227647	j,y,z		EST, Highly similar to T14106 probable GTPase-activating protein SPA-1 - rat [R.norvegicus], ESTs, Highly similar to T14106 probable GTPase-activating protein SPA-1 - rat [R.norvegicus], KIAA0440 protein, RIKEN cDNA 2610511M17 gene, expressed sequence AW213287, signal-induced proliferation associated gene 1 hypothetical protein, clone 1-53, small inducible cytokine subfamily D (Cys-X3 Cys), member 1 (fractalkine, neurotactin), small inducible cytokine subfamily D, 1
1178	22838	AI227667	aa		Substantilly D, 1
1179	6765	AI227761	i,General		KIAA0665 gene product, KIAA1821 protein
	10,00	1, 1,22,1,01	i, ochorai	[Protoni

Son	TARI B R	e MINMANI	പ്രയ്യാ യുപ്പട്ട ഗു	SKIOTE/TEOKI		Atty, Docket No. 44921-5089W0
						Dos. No. 1798397.1
Ilinkad 1, hypothetical protein	Sog. (D No:	ldentifier	ConBank Ace./ Ref. Seq. ID No.			Homologous Cluster Name
1182 23898 Al229987 d peptidy(glycine alpha-amidating amidating monoxygenase peptidy(glycine alpha-amidating monoxygenase Store peptidy(glycine alpha-amidating monoxygenase peptidy(glycine alpha-amidating monoxygenase peptidy(glycine alpha-amidating monoxygenase peptidy(glycine alpha-amidating monoxygenase peptidy(glycine alpha-amidating monoxygenase peptidy(glycine alpha-amidating monoxygenase peptidy(glycine alpha-amidating monoxygenase peptidy(glycine alpha-amidating monoxygenase peptidy(glycine alpha-amidating monoxygenase peptidy(glycine alpha-amidation gestycle alpha peptidy(glycine alpha-amidation peptidy(glycine alpha-amidation peptidy(glycine alpha-amidation peptidy(gl				General		FLJ10097, nerve growth factor receptor (TNFRSF16) associated
183						
1183 1651 Al228068 n,w amidaling monooxygenase mon	1102	23090	A1221901	u	pentidylalycine alpha-	nentidylalycine alpha-amidating
1184 14237 Al228128 e	1183	1651	A1228068	n w		
1185					arriadaring monecky genese	Interior ygoridoo
1186						C21I_HUMAN PUTATIVE PROTEIN C21ORF18 [H.sapiens], chromosome
1187 22915 Al228299 r		- 22				21 open reading frame 18
1188 8917 Al228313 General				0		
1189				Gonoral		
1191						
1191 1192 13730 13745 13730 13745 13730 13745 13730 13745 13730 13745 13730 13745 13730 13745 13730 13745 13730 13745 13730 13745 13730 13745 13730 13745 13730 13745 13730 13745 13730 13745 13730 13745 13730 13745 13730 13745						
ESTS, Highly similar to S70642 ubiquith ligase Nedd4 - rat [R.norvegicus], ESTS, Moderately similar to S70642 ubiquith ligase Nedd4 - rat [R.norvegicus], ESTS, Moderately similar to S70642 ubiquith ligase Nedd4 - rat [R.norvegicus], ESTS, Weakly similar to S70642 ubiquith ligase Nedd4 - rat [R.norvegicus], ESTS, Weakly similar to NED4 MOUSE NED0.4 PTOETIN [M.musculus], RIKEN cDNA 2600016C11 gene, neural precursor cell expressed, developmentally down-regulated qn evelopmentally down-regulated gen 4a						
1193 13745 Al228494 b,cc hypothetical protein MGC4614, membrane component, chromosome 17, surface marker 2 (ovarian carcinoma antigen CA125), next to the Broat, oxidative stress induced like, sequestosome 1 1194	1192	13730	A1228356	a		ubiquitin ligase Nedd4 - rat [R.norvegicus], ESTs, Moderately similar to S70642 ubiquitin ligase Nedd4 - rat [R.norvegicus], ESTs, Weakly similar to NED4 MOUSE NEDD-4 PROTEIN [M.musculus], RIKEN cDNA 2600016C11 gene, neural precursor cell expressed, developmentally down-regulated 4, neural precursor cell expressed, developmentally down-regulated gene
hypothetical protein MGC4614, membrane component, chromosome						
1195						membrane component, chromosome 17, surface marker 2 (ovarian carcinoma antigen CA125), next to the Brca1, oxidative stress induced like,
1197 11605 Al228682 e 1198 13203 Al228728 r DEAD/H (Asp-Glu-Ala-Asp/His) box binding protein 1, Protein inhibitor of activated STAT X, protein inhibitor of activated STAT protein PIASy, protei inhibitor of activated STAT activated STAT activated STAT protein PIASy, protei inhibitor of activated STAT protein PIASy, protein inhibitor of activated STAT activated S	1195	16053	Al228596	CC		
1198 13203 Al228728 r DEAD/H (Asp-Glu-Ala-Asp/His) box binding protein 1, Protein inhibitor of activated STAT X, protein inhibitor of activated STAT protein PlASy, protei inhibitor of activated STAT protein PlASy, protei inhibitor of activated STAT3 1200	1196	3557	AI228672	е		
DEAD/H (Asp-Glu-Ala-Asp/His) box binding protein 1, Protein inhibitor of activated STAT x, protein inhibitor of activated STAT protein inhibitor of activated STAT protein PIASy, protein inhibitor of activated STAT protein PIASy, protein inhibitor of activated STAT protein PIASy, protein inhibitor of activated STAT protein PIASy, protein inhibitor of activated STAT protein PIASy, protein protein 28, vesicle-associated membrane protein 28, vesicle-associated membrane protein 1, vesicle-associated membrane protein 1, vesicle-associated membrane protein 1, vesicle-associated membrane protein 1, vesicle-associated membrane protein 1, vesicle-associated membrane protein 1, vesicle-associated membrane protein 1, vesicle-associated membrane protein 1, vesicle-associated membrane protein 1, vesicle-associated membrane protein 5, vesicle-associated membrane protein 5, vesicle-associated membrane protein 8, vesicle-associated membrane protein 8, vesicle-associated membrane protein 8, vesicle-associated membrane protein 8, vesicle-associated membrane protein 8, vesicle-associated membrane protein 8, vesicle-associated membrane protein 8, vesicle-associated membrane protein 8, vesicle-associated membrane protein 8, vesicle-associated membrane protein 8, vesicle-associated membrane protein 8, vesicle-associated membrane protein 8, vesicle-associated membrane protein 8, vesicle-associated membrane protein 8, vesicle-associated membrane protein 8, vesicle-associated membrane protein 8, vesicle-associated membrane protein 8, vesicle-associated membrane protein 9, vesicle-associated membrane protein 9, vesicle-associated membrane protein 9, vesicle-associated membrane protein 9, vesicle-associated membrane protein 9, vesicle-associated membrane protein 9, vesicle-associated membrane protein 9, vesicle-associated membrane protein 9, vesicle-associated membrane protein 9, vesicle-associated membrane protein 9, vesicle-associated membrane protein 9, vesicle-associated membrane protein 9, vesicle-associated membrane protein 9, vesic			Al228682	е		
1201 8235 Al229154 k Rattus norvegicus mRNA for vesicle associated membrane protein 2B, vesicle-associated membrane protein 1, vesicle-associated membrane protein 1 (synaptobrevin 1), vesicle-associated membrane protein 1, vesicle-associated membrane protein 1, vesicle-associated membrane protein 1 (synaptobrevin 2), vesicle-associated membrane protein 1 (synaptobrevin 1), vesicle-associated membrane protein 5, vesicle-associated membrane protein 5, vesicle-associated membrane protein 6, vesicle-associated membrane protein 8, vesicle-associated membrane protein 8, vesicle-associated membrane protein 8 (endobrevin) 1202 16203 Al229196 r protein 2 (synaptobrevin 2) (endobrevin)	1199	13771	Al228848	g		binding protein 1, Protein inhibitor of activated STAT X, protein inhibitor of activated STAT protein PIASy, protein
Rattus norvegicus mRNA for vesicle associated membrane protein 2B, vesicle-associated membrane protein 1, vesicle-associated membrane protein 1 (synaptobrevin 1), vesicle-associated membrane protein 2, vesicle-associated membrane protein 2 (synaptobrevin 2), vesicle-associated membrane protein 1 (synaptobrevin 1), vesicle-associated membrane protein 5, vesicle-associated membrane protein 5 (myobrevin), vesicle-associated membrane protein 8, vesicle-associated membrane protein 8 (endobrevin) 1202 16203 Al229196 r protein 2 (synaptobrevin 2) (endobrevin)				<u> </u>		
1203 13826 Al229304 a				K	membrane protein 1, vesicle-associated membrane protein 1 (synaptobrevin 1), vesicle- associated membrane protein 2, vesicle- associated membrane	associated membrane protein 2B, vesicle-associated membrane protein 1, vesicle-associated membrane protein 1 (synaptobrevin 1), vesicle-associated membrane protein 2, vesicle-associated membrane protein 2 (synaptobrevin 2), vesicle-associated membrane protein 5, vesicle-associated membrane protein 5 (myobrevin), vesicle-associated membrane protein 8, vesicle-associated membrane protein 8
				r	protein 2 (synaptobrevin 2)	(endobrevin)

TABLES	e HUMAN	HOMOLOGUE AN	SKOLLVALLOKI		Atty, Docket No. 449211-5089W0
Seq. [D	ldentijer	CenBank Ace./ + Ref. Seq. ID No.;	Medal Code	Homologous Cana Nama	Doc. No. 1793397.1 Homologous Gluster Nama
1205	4640	Al229404	x.aa	wems	maniprogram arcettar intaling
1206	23563	AI229421	I		MAP kinase-activated protein kinase 2, mitogen-activated protein kinase- activated protein kinase 3 NADH dehydrogenase (ubiquinone) 1
1207	15426	AI229497	s		beta subcomplex, 10 (22kD, PDSW)
1208	15193	AI229508	bb		
1209	19243	AI229638	×		thymidine kinase 1, thymidine kinase 1, soluble
1210	23078	AI229647	p		I, soluble
1211	3099	A1229680	o		NADH dehydrogenase (ubiquinone) Fe- S protein 3 (30kD) (NADH-coenzyme Q reductase)
1212	19508	AI229698	bb		
1213	13977	Al229707	×		EST, Moderately similar to I38369 beta tubulin [H.sapiens], EST, Weakly similar to I38369 beta-tubulin [H.sapiens], EST, Weakly similar to TUBULIN BETA-5 CHAIN [M.musculus], ESTs, Highly similar to A25113 tubulin beta chain 15 - rat [R.norvegicus], FK506-binding protein 1A (12kD), RIKEN cDNA 2310061K05 gene, RIKEN cDNA 2410129E14 gene, tubulin, beta 2, tubulin, beta 5, tubulin, beta polypeptide
				,	quinolinate phosphoribosyltransferase
1214	23983	Al229708	V		(nicotinate-nucleotide pyrophosphorylase (carboxylating))
1215	2688	Al229793	e		pyrophosphorylase (carboxylating))
1216	13874	Al229832	9		
1217 1218	12587 20591	Al229979 Al229993	General I,m		ESTs, Weakly similar to MOT2 MOUSE MONOCARBOXYLATE TRANSPORTER 2 [M.musculus], monocarboxylate transporter 4, solute carrier 16 (monocarboxylic acid transporters), member 8, solute carrier family 16 (monocarboxylic acid transporters), member 3, solute carrier family 16 (monocarboxylic acid transporters), member 7, solute carrier family 16 (monocarboxylic acid transporters), member 7, solute carrier family 16 (monocarboxylic acid transporters), member 8
1219	24042	AI230002	a.b.d.General		
1220	13880	Al230042	u		EST, Highly similar to CCAA MOUSE VOLTAGE-DEPENDENT P/Q-TYPE CALCIUM CHANNEL ALPHA-1A SUBUNIT [M.musculus], ESTs, Weakly similar to CCAA MOUSE VOLTAGE-DEPENDENT P/Q-TYPE CALCIUM CHANNEL ALPHA-1A SUBUNIT [M.musculus], Homo sapiens mRNA; cDNA DKFZp434M0223 (from clone DKFZp434M0223); partial cds, Mus musculus calcium channel mRNA, complete cds, calcium channel, P/Q type, alpha 1A, calcium channel, voltage-dependent, alpha 1G subunit, two-pore channel 1, homolog
1221	17672	A1230074	d		NADH dehydrogenase (ubiquinone) 1 alpha subcomplex, 1, NADH dehydrogenase (ubiquinone) 1 alpha subcomplex, 1 (7.5kD, MWFE)

TABLE	: HUMAN	HOMOROGUE AR	. EKIORIATOKI	Atom A court	Atty. Docket No. 44921-5039W0
Seq. ID	10000000	ConBenk Acc.	Model Code	Konologovs Cone	Dog. No. 1793397.1
NO ₅	Identifier	Rei. Seg. 10 No.	Model Gode	Name ::	Homologous Cluster Name ESTs, Highly similar to E2BE RAT TRANSLATION INITIATION FACTOR
					EIF-2B EPSILON SUBUNIT [R.norvegicus], HSPC028 protein,
					RIKEN cDNA 1200015E15 gene, Rattus norvegicus initiation factor eIF-
					2Be mRNA, complete cds, basic leucine-zipper protein BZAP45,
					eukaryotic translation initiation factor 2B, subunit 5 (epsilon, 82kD),
1222	3652	AI230113	General		expressed sequence C81315 EST, Weakly similar to ENPL HUMAN
					ENDOPLASMIN PRECURSOR
					[H.sapiens], ESTs, Highly similar to HS9B RAT HEAT SHOCK PROTEIN
					HSP 90-BETA [R.norvegicus], ESTs, Weakly similar to ENPL_HUMAN
					ENDOPLASMIN PRECURSOR [H.sapiens], Homo sapiens mRNA;
					cDNA DKFZp564F053 (from clone DKFZp564F053), RIKEN cDNA
					1810014B01 gene, RIKEN cDNA 2410002K23 gene, expressed
					sequence C81438, heat shock 90kD protein 1, beta, heat shock protein, 84
1223	18650	AI230121	aa		kDa 1, tumor rejection antigen (gp96) 1, tumor rejection antigen gp96
	5				ESTs, Moderately similar to CHD3_HUMAN CHROMODOMAIN HELICASE-DNA-BINDING PROTEIN
1224	13025	Al230173	С		3 [H.sapiens], chromodomain helicase DNA binding protein 3
				·	ESTs, Highly similar to A47327 selenoprotein P precursor [H.sapiens],
1225 1226	4280 18528	AI230247 AI230284	z General	selenoprotein P, plasma, 1	selenoprotein P, plasma, 1
1227	7084	AI230362	р		Homo sapiens, clone IMAGE:3845253, mRNA, partial cds
1228 1229	20895 12961	AI230549 AI230554	b,n General		
1229	12901	A1230554	General	_	EST, Moderately similar to GALECTIN-
					1 [R.norvegicus], Human HL14 gene encoding beta-galactoside-binding
					lectin, 3' end, clone 2, RIKEN cDNA 2200008F12 gene, beta-galactoside-
					binding lectin, lectin, galactose binding, soluble 1, lectin, galactoside-
					binding, soluble, 1 (galectin 1), lectin, galactoside-binding, soluble, 2
1230	15636	AI230616	r		(galectin 2)
1231	4121	AI230647	lj,m		DNA segment, Chr 17, ERATO Doi
1232	14388	AI230702	General		441, expressed, hematological and neurological expressed 1
1233	18529	Al230716	x,General		EST, Weakly similar to JW0105
					synaptojanin 2 alpha protein - mouse [M.musculus], ESTs, Weakly similar to
					2204390A synaptojanin [R.norvegicus], ESTs, Weakly similar
					to JW0105 synaptojanin 2 alpha
				i	protein - mouse [M.musculus], expressed sequence AA675315,
	10046	41000704			inositol polyphosphate-5-phosphatase, 75kD, putative phosphatase,
1234 1235	13618 8304	Al230724 Al230746	General cc		suppressor of actin 1, synaptojanin 2

TABLE	HUMAN	HOMOLOGUEA	· SKOTATOKI	Transfer (27 Fileson)	Atty. Docket No. 44921-5089W0
Seq. [D		Genbenk Acc.		Homologous Cene	Doc. No. 1793397.1
No.	ldeniiier	Ref. Seq. ID No.	Model Gode	Name : : :	Homologous Cluster Name
1236	4731	Al230773	е		
1237	14430	Al230798	c,k,x		
1238	16627	Al230822	bb		Alg5, S. cerevisiae, homolog of, dolichyl-phosphate mannosyltransferase polypeptide 1, catalytic subunit
1239 1240	3125 633	AI231028 AI231127	General k		DNA segment, Chr 10, ERATO Doi 398, expressed, ESTs, Weakly similar to PTNL RAT PROTEIN TYROSINE PHOSPHATASE, NON-RECEPTOR TYPE 21 [R.norvegicus], Homo sapiens, Similar to erythrocyte membrane protein band 4.1-like 3, clone MGC:12343 IMAGE:4044866, mRNA, complete cds, Mus musculus adult male pituitary gland cDNA, RIKEN full-length enriched library, clone:5330430110, full insert sequence, Rattus norvegicus protein tyrosine phosphatase 2E (PTP2E) mRNA, complete cds, erythrocyte membrane protein band 4.1 (elliptocytosis 1, RH-linked), erythrocyte membrane protein band 4.1-like 1, erythrocyte protein band 4.1-like 3, erythrocyte protein band 4.1-like 3, protein tyrosine phosphatase, non-receptor type 21
1241	20846	Al231140	P		EST, Highly similar to 60S RIBOSOMAL PROTEIN L23A [R.norvegicus], ESTs, Highly similar to RL2B_HUMAN 60S RIBOSOMAL PROTEIN L23A [H.sapiens], ribosomal protein L23a
1242	6743	Al231219	ld .		protein c23a
1244	26292	Ai231391	k		
1245	12343	Al231433	w	· · · · · · · · · · · · · · · · · · ·	
1246	7337	Al231465	aa		
1247	16321	AI231506	General	· · · · · · · · · · · · · · · · · · ·	
1248	8004 15171	Al231532 Al231792	j,l g		ESTs, Moderately similar to Z183_HUMAN ZINC FINGER PROTEIN 183 [H.sapiens], zinc finger protein 183 (RING finger, C3HC4 type) BCL2-associated athanogene 3, Bcl2- associated athanogene 3, RIKEN cDNA 1600025G07 gene, RIKEN cDNA 2410112I15 gene
1250	6193	Al231797	1		
1252	14227	AI231999	u		RIKEN cDNA 2810411G23 gene, tumor protein D52-like 1 EST, Moderately similar to EF1D_HUMAN ELONGATION FACTOR 1-DELTA [H.sapiens], ESTs, Weakly similar to EF1D_HUMAN ELONGATION FACTOR 1-DELTA [H.sapiens], eukaryotic translation elongation factor 1 delta (guanine nucleotide exchange protein),
	124504	AI232006	w,y,bb	Ī	hypothetical protein FLJ20897
1253 1254	24501	A1232000	g,q,z,cc,General		Inypothetical protein i Edzoosi

	TABLES	HUMAN	HOMOTOGNE W	INOTATIONS		Atty. Docket No. 44921-5039WO
					loo	<u> </u>
Transistion Initiation Factor Eff1, Nmr, 29 Structures (Hapier, RiKEN cDNA 1500010M14 gene, RikEN cDNA 310001M14 gene, putative translation initiation factor, suppressor of initiator codor mutation related sequence 1 (\$5. cerevisiae) 1255				Model.Code	Name	Homologous Gluster Name
1256						ESTs, Highly similar to Human Translation Initiation Factor Eif1, Nmr, 29 Structures [H.sapiens], RIKEN cDNA 1500010M16 gene, RIKEN cDNA 3110001N14 gene, putative translation initiation factor, suppressor of initiator codon mutations, related
1525						sequence 1 (5. cerevisiae)
1259 21326 132274 Lm						
1259						
1261 2913 Al232272 0		23125	AI232266			
1263	1260	2085	Ál232270	bb		
1263	1261	2913	Al232272	0		
1264	1262	14304	Al232281	g		
1264 15122	1263	15955	Al232294	u,bb,General		
Durinergic receptor P2X, ligand-gated ion channel 4, purinergic receptor P2X, ligand-gated ion channel, 4		45422				sapiens chromosome 19, cosmid R31343, RIKEN cDNA 1700065B19 gene, RIKEN cDNA 5730408C10 gene, likely ortholog of mouse g1- related zinc finger protein, ring finger
Durinergic receptor PZX, ligand-gated ion channel 4, purinergic receptor PZX, ligand-gated ion channel, 1, purinergic receptor PZX, ligand-gated ion channel, 1, purinergic receptor PZX, ligand-gated ion channel, 4 ligand-gated ion cha	1264	15122	A1232303	У		
1267 24321 Al232340 O factor 1 stromal cell derived factor 1 stromal cell-derived factor 1 stromal cell-derived factor 1 stromal cell-derived factor 1 cel				у	ligand-gated ion channel 4, purinergic receptor P2X,	ion channel 4, purinergic receptor P2X, ligand-gated ion channel, 1, purinergic receptor P2X, ligand-gated ion
1267 24321 Al232340 O factor 1 Stromal cell-derived factor 1, Stromal cell-derived factor 1, Stromal cell-derived factor 1	1266	15246	A1232332	t,u		
1268	4007	04004	A 10000 40		1, stromal cell-derived	
1270					factor 1	cell-derived factor 1
1270 19287 Al232379 f receptor, alpha polypeptide, platelet derived growth factor receptor, alpha polypeptide, platelet derived growth factor receptor, alpha polypeptide						
Tactor receptor, alpha polypeptide, platelet-derived growth factor receptor, alpha polypeptide, platelet-derived growth factor receptor, alpha polypeptide Flavin-containing monooxygenase 1, flavin containing monooxygenase 1, flavin containing monooxygenase 2, flavin containing monooxygenase 3, flavin containing monooxygenase 3, flavin containing monooxygenase 3, flavin containing monooxygenase 4, hypothetical protein PRO1257 1272	1269	11411	A1232346	In .		
Section Figure	1270	19287	Al232379	f	factor receptor, alpha polypeptide, platelet- derived growth factor	
ESTs, Moderately similar to A27340 complement C7 precursor [H.sapiens], complement C7 precursor [H.sapiens], complement component 7 1274 11157 Al232494 cc DnaJ (Hsp40) homolog, subfamily B, member 3, DnaJ (Hsp40) homolog, subfamily B, member 6, ESTs, Moderately similar to HSJ2_HUMAN DNAJ PROTEIN HOMOLOG 2 [H.sapiens], ESTs, Weakly similar to HSJ2_HUMAN DNAJ PROTEIN HOMOLOG 2 [H.sapiens], Homo sapiens, clone IMAGE:3930327, mRNA, partial cds, RIKEN cDNA 2810451A06 gene, expressed sequence AU020082 1275 8709 Al232534 o sequence AU020082 1276 20350 Al232552 j,v,y 1277 14069 Al232631 e						flavin containing monooxygenase 1, flavin containing monooxygenase 2, flavin containing monooxygenase 3, flavin containing monooxygenase 4,
1273 5572 Al232490 i,t complement C7 precursor [H.sapiens], complement component 7	1212	14001	MIZ32409	1,111		ESTs Madorataly similar to A27240
1273 5572 Al232490 i,t complement component 7 1274 11157 Al232494 cc DnaJ (Hsp40) homolog, subfamily B, member 3, DnaJ (Hsp40) homolog, subfamily B, member 6, ESTs, Moderately similar to HSJ2_HUMAN DNAJ PROTEIN HOMOLOG 2 [H.sapiens], ESTs, Weakly similar to HSJ2_HUMAN DNAJ PROTEIN HOMOLOG 2 [H.sapiens], Homo sapiens, clone IMAGE:3930327, mRNA, partial cds, RIKEN cDNA 2810451A06 gene, expressed sequence AU020082 1275 8709 Al232534 o sequence AU020082 1276 20350 Al232552 j.v.y						
1274 11157 Al232494 cc DnaJ (Hsp40) homolog, subfamily B, member 3, DnaJ (Hsp40) homolog, subfamily B, member 3, DnaJ (Hsp40) homolog, subfamily B, member 6, ESTs, Moderately similar to HSJ2_HUMAN DNAJ PROTEIN HOMOLOG 2 [H. sapiens], ESTs, Weakly similar to HSJ2_HUMAN DNAJ PROTEIN HOMOLOG 2 [H. sapiens], Homo sapiens, clone IMAGE:3930327, mRNA, partial cds, RIKEN cDNA 2810451A06 gene, expressed sequence AU020082 1275 8709 Al232534 o sequence AU020082 1276 20350 Al232552 j.v.y 1277 14069 Al232631 e	1273	5572	A1232490	i t		
DnaJ (Hsp40) homolog, subfamily B, member 3, DnaJ (Hsp40) homolog, subfamily B, member 3, DnaJ (Hsp40) homolog, subfamily B, member 6, ESTs, Moderately similar to HSJ2_HUMAN DNAJ PROTEIN HOMOLOG 2 [H.sapiens], ESTs, Weakly similar to HSJ2_HUMAN DNAJ PROTEIN HOMOLOG 2 [H.sapiens], Homo sapiens, clone IMAGE:3930327, mRNA, partial cds, RIKEN cDNA 2810451A06 gene, expressed sequence AU020082 j.v.yy 1277 14069 AI232531 e						complement component /
1277 14069 Al232631 e	1275	8709	Al232534	o		member 3, DnaJ (Hsp40) homolog, subfamily B, member 6, ESTs, Moderately similar to HSJ2_HUMAN DNAJ PROTEIN HOMOLOG 2 [H.sapiens], ESTs, Weakly similar to HSJ2_HUMAN DNAJ PROTEIN HOMOLOG 2 [H.sapiens], Homo sapiens, clone IMAGE:3930327, mRNA, partial cds, RIKEN cDNA 2810451A06 gene, expressed
			Al232631			
	1277	4440	Al232643	w	-	

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		KOWOTOGNE W			Aity, Docket No. 44921-5039W0 Doc. No. 1798397.1
839. ID	ldeniiiier	Cenbank Ace./ Ref. Seq. ID No.	Model Code	Homologous Cene Name	Homologous Cluster Name
1279	17695	AI232784	е		2,4-dienoyl CoA reductase 1, mitochondrial, 2,4-dienoyl CoA reductase 2, peroxisomal, ESTs, Weakly similar to S11021 2,4-dienoyl-CoA reductase [R.norvegicus], FabG (beta-ketoacyl-[acyl-carrier-protein] reductase, E coli) like, H2-K region expressed gene 6, Homo sapiens AS10 protein mRNA, partial cds, Mus musculus, clone MGC:6971 IMAGE:3154595, mRNA, complete cds, RIKEN cDNA 0610039E24 gene, RIKEN cDNA 1810026B04 gene, carbonyl reductase, oxidoreductase UCPA, peroxisomal trans 2-enoyl CoA reductase; putative short chain alcohol dehydrogenase
1280	15796	Al232874	v		
1281	12467	A1222084	General		ESTs, Highly similar to I49636 DNA-binding protein - mouse [M.musculus], ESTs, Moderately similar to I49636 DNA-binding protein - mouse [M.musculus], ESTs, Weakly similar to OZF_HUMAN ZINC FINGER PROTEIN OZF [H.sapiens], RIKEN cDNA 2310011F05 gene, pancreas zinc finger protein, zinc finger protein 260, zinc finger protein 36 (KOX 18), zinc finger protein 63, zinc finger
1282	12873	AI232984	[]		protein 97
1283 1284	5355 18794	AI233031 AI233121	r		
1285	3823	Al233147	b,g,General		DNA segment, Chr 17, human D6S81E 1, ESTs, Highly similar to S33681 translation initiation factor eIF- 4A.I [H.sapiens], HLA-B associated transcript 1, Human clone 23933 mRNA sequence, eukaryotic translation initiation factor 4A, isoform 1, mitochondrial DEAD-box polypeptide 28, nuclear RNA helicase, DECD variant of DEAD box family
1286	11967	Al233155	c,k,General		DECD Variant of DEAD box family
1287	11561	Al233182	d		
1288	3471	Al233183	g		putative receptor protein
	21948	Al233191	i		
1290	13598	AI233194	g,p,y		
1291	15552	Al233195	у		
					EST, Moderately similar to EGFR_HUMAN EPIDERMAL GROWTH FACTOR RECEPTOR PRECURSOR [H.sapiens], Epidermal growth factor receptor, formerly avian erythroblastic leukemia viral (v-erbB) oncogene homolog (Erbb1), avian erythroblastosis oncogene B 3, epidermal growth factor receptor, epidermal growth factor receptor (avian erythroblastic leukemia viral (v-
	17907	AI233224	bb		erb-b) oncogene homolog)
1293	14111	Al233269	СС		
1294	12894	Al233365	d		
1295	7161	A1233407	General		
1296 1297	15906 14120	Al233425	q		
		AI233433 AI233468	d a d		
1290	14033	NIZ33400	a,d		

TABLE 3	: HUMAN	HOWOLOGAE W	NOTATIONS .		Atty. Docket No. 44921-5089W0
Seq. (D		ConBank Ace.		SHOUND STAND	<u> </u>
		Rei Seq. ID No.	Model Code	Name :	Homologova Cluater Name
1299	3075	AI233494	u,aa		oxidase (cytochrome c) assembly 1- like
1300	6046	AI233530	General		
					EST, Moderately similar to S56108 26S proteasome regulatory complex chain p31 [H.sapiens], proteasome (prosome, macropain) 26S subunit,
1301	18900	AI233570	General		non-ATPase, 8 Homo sapiens, clone MGC:14993
1302	7888	Al233583	General		IMAGE:3613406, mRNA, complete cds, arginyl-tRNA synthetase
1303	16709	AI233602	General	adenosine kinase	adenosine kinase, expressed sequence Al255373
1304	5163	AI233712	у		protein phosphatase 1D magnesium- dependent, delta isoform
1305	7243	Al233717	General		coatomer protein complex, subunit alpha, embryonic ectoderm development
1306	3816	Al233729	g		Homo sapiens cDNA FLJ10203 fis, clone HEMBA1004930, moderately similar to 26S PROTEASOME SUBUNIT S5B, proteasome (prosome, macropain) 26S subunit, non-ATPase, 5
1307	13023	Al233740	d,h,General		Aldehyde reductase 1 (low Km aldose reductase) (5.8 kb Pstl fragment, probably the functional gene), EST, Highly similar to Aldose Reductase (E.C.1.1.1.21) [H.sapiens], ESTs, Moderately similar to ALDOSE REDUCTASE [R.norvegicus], ESTs, Moderately similar to ALDOSE REDUCTASE-RELATED PROTEIN 2 [M.musculus], Homo sapiens, Similar to RIKEN cDNA 1110018J12 gene, clone IMAGE:3865164, mRNA, partial cds, RIKEN cDNA 2310005E10 gene, aldo-keto reductase family 1, member A4 (aldehyde reductase), aldo-keto reductase family 1, member B1 (aldose reductase), aldo-keto reductase family 1, member B10 (aldose reductase), in member B3 (aldose reductase), fibroblast growth factor regulated protein
1308	14871	Al233743	g g		
1309	7469	Al233767	cc		Golgi-associated, gamma-adaptin ear containing, ARF-binding protein 2
1310	7804	Al233767	b		containing, Arti -bilding protein Z
1311	13563	AI233773	e		
1312	2154	Al233818	k,cc	A kinase (PRKA) anchor protein (gravin) 12	
1313	16616	Al234079	h	15	
1314	13393	Al234100	a,d,General		EST, Moderately similar to CYSR RAT CYSTEINE-RICH PROTEIN 1 [R.norvegicus], ESTs, Weakly similar to CYSR RAT CYSTEINE-RICH PROTEIN 1 [R.norvegicus], ESTs, Weakly similar to S12658 cysteine-rich protein [H.sapiens], cysteine and glycine-rich protein 1, cysteine rich protein, cysteine-rich protein 2, cysteine-rich protein 3, epithelial protein lost in neoplasm beta, thymus LIM protein
1315	7071	AI234162	r		

TABLES	RUMAN	HOMOLOGUE AN	INOTATIONS		Auy. Docket No. 44921-5039WC Doc. No. 1793397.1
809. [D	ldeniliter	Cenbank Ace.// Ref. Seq. (D No.	Model Gode	Hemelogous Gene Name	Homologous Cluster Namo
1316	14677	AI234620	General		
1317	4443	AI234629	m		
1318	22453	AI234678	b		
1319	23964	Al234748	t,General		
1320	19581	AI234753	f		
1321 1322	22152 18942	AI234822 AI234865	o,General	RAS, dexamethasone- induced 1	Harvey rat sarcoma oncogene, subgroup R, RAP1B, member of RAS oncogene family, RAP2B, member of RAS oncogene family, RIKEN cDNA 2610016H24 gene, RIKEN cDNA 4021402C18 gene, RIKEN cDNA 4930526B11 gene, RIKEN cDNA 5830461H18 gene, expressed sequence Al573426, rap2A-like protein, ras-related protein
1322	18942	A1234800			ATDess III Assessment I have seen
1323	22662	Al234939	aa		ATPase, H+ transporting, lysosomal (vacuolar proton pump), subunit 1, EST, Weakly similar to 154197 hypothetical protein [H.sapiens], ESTs, Weakly similar to VAS1_RAT VACUOLAR ATP SYNTHASE SUBUNIT S1 PRECURSOR (V-ATPASE S1 SUBUNIT) (V-ATPASE S1 SUBUNIT) (V-ATPASE S1 SUBUNIT) (C7-1 PROTEIN) [R.norvegicus], Homo sapiens cDNA FLJ12563 fis, clone NT2RM4000820, weakly similar to VACUOLAR ATP SYNTHASE SUBUNIT AC45 PRECURSOR (EC 3.6.1.34)
4004	0075	A1005047			CB80_HUMAN 80 KDA NUCLEAR CAP BINDING PROTEIN [H.sapiens], Homo sapiens cDNA FLJ11599 fis, clone HEMBA1003879, nuclear cap
1324	3875	AI235047	o,General		binding protein subunit 1, 80kD
1326	14906	AI235135	g		ATP-binding cassette, sub-family F (GCN20), member 1, ATP-binding cassette, sub-family F (GCN20), member 2, hypothetical protein FLJ11198
1327	14718	AI235210	е		FOT Moderately similar to TIMA DAT
1328 1329	15004 6632	AI235224 AI235277	b,General		EST, Moderately similar to TIM1 RAT METALLOPROTEINASE INHIBITOR 1 PRECURSOR [R.norvegicus], EST, Weakly similar to TIM1 RAT METALLOPROTEINASE INHIBITOR 1 PRECURSOR [R.norvegicus], tissue inhibitor of metalloproteinase, tissue inhibitor of metalloproteinase 1 (erythroid potentiating activity, collagenase inhibitor)
1329	0032	A1233211	V	 	ESTa Highly similar to ASSESS T sell
1330 1331	14722 1462	Al235284 Al235585	x,z u,General		ESTs, Highly similar to A60592 T-cell surface glycoprotein E2 precursor [H.sapiens], Homo sapiens cDNA FLJ13471 fis, clone PLACE1003566, antigen identified by monoclonal antibodies 12E7, F21 and O13, hypothetical protein DKFZp761H2024
1332	21061	Al235631	I,m		
1334	121001	MIZOUO I	[1,111	l	

TABLES	g IHUMAN	HOWO TO GATE WY	NOTATIONS		Ally. Docket No. 44921-5089W0 Doc. No. 1793897.1
Seq. (D No. :	ldentiller	GenBank Acc./ Ref. Seq. ID No.) 	Homologous Gana Namo	Homologovs Gluster Name
4222	44665	Algaecac		MAD (mothers against decapentaplegic, Drosophila) homolog 4, MAD homolog 4	MAD (mothers against decapentaplegic, Drosophila) homolog
1333	14665	AI235646	M Conord	(Drosophila)	4
1334 1335	19940 5698	Al235689 Al235692	General u		
1333	3030	A1203032	<u>u</u>		ESTs, Weakly similar to LDVR RAT VERY LOW-DENSITY LIPOPROTEIN RECEPTOR PRECURSOR [R.norvegicus], RIKEN cDNA 4933405111 gene, Very low density lipoprotein receptor, nidogen 2, secreted modular calcium-binding protein 1, secreted modular calcium-binding protein 2, very low density
1336	23745	Al235732	k		lipoprotein receptor ESTs, Highly similar to A56716 aromatic ester hydrolase [H.sapiens], biphenyl hydrolase-like (serine hydrolase; breast epithelial mucin-
1337	11164	AI235739	General		associated antigen)
1338	5212	AI235745	d		
1339	14768	Al235912	h		
1340	14776	AI235950	m		
1341	3091	AI236027	n,General		
1342	14861	AI236045	<u>r</u>		
1343	14862 16943	Al236048 Al236097	e		integral membrane protein 2 B, integral membrane protein 2B, integral membrane protein 3
1345	8336	Al236101	p I		memorane protein 3
1346	23230	Al236146	v		
1347	22855	Al236150	e		Down syndrome critical region gene 5
1348	14594	AI236152	i		Dewit Syndrome Childer region gene 5
1349	18406	Al236168	r	-	
1350	15051	Al236332	General		ESTs, Weakly similar to ATDA_HUMAN DIAMINE ACETYLTRANSFERASE [H.sapiens], RIKEN cDNA 2610016A03 gene, RIKEN cDNA 4930404K22 gene, spermidine/spermine N1-acetyl transferase, spermidine/spermine N1- acetyltransferase
1351	19298	AI236338	bb		ESTs, Highly similar to NHPX RAT NHP2/RS6 FAMILY PROTEIN YEL026W HOMOLOG [R.norvegicus], RIKEN cDNA 2410130M07 gene, nonhistone chromosome protein 2 (S. cerevisiae)-like 1, nucleolar protein family A, member 2 (H/ACA small nucleolar RNPs), sperm specific antigen 1

		HOMOFÓGAÉ W			Atty. Docket No. 44921-5089W0 Doc. No. 1798897.1
Seq. ID No.	ldeniiiier	Confort Ace./ Ref. Seq. ID No.	Model Code	Homologous Gene Name	Homologous Cluster Name
					ELAV (embryonic lethal, abnormal vision, Drosophila)-like 4 (Hu antigen D), EST, Highly similar to 2201474A inducible poly(A)-binding protein [H.sapiens], EST, Weakly similar to 2201474A inducible poly(A)-binding protein [H.sapiens], ESTs, Weakly similar to 2201474A inducible poly(A)-binding protein [H.sapiens], ESTs, Highly similar to HUD RAT PARANEOPLASTIC ENCEPHALOMYELITIS ANTIGEN HUD HOMOLOG [R.norvegicus], ESTs, Moderately similar to HUD RAT PARANEOPLASTIC ENCEPHALOMYELITIS ANTIGEN HUD HOMOLOG [R.norvegicus], ESTs, Moderately similar to POLYADENYLATE-BINDING PROTEIN 1 [M.musculus], Mus musculus adult male testis cDNA, RIKEN full-length enriched library, clone:4933407N23, full insert sequence, RIKEN cDNA 4932702K14 gene, poly A binding protein, cytoplasmic 1, poly(A)-binding protein, cytoplasmic 4 (inducible form), siah binding protein 1; FBP interacting repressor; pyrimidine tract binding splicing factor; Ro ribonucleoprotein-
1352	10667	AI236366	b		binding protein 1
1353	10774	AI236397	f		
1354	9407	AI236402	aa		
1355	26335	Al236460_	General		
1356	17950	AI236590	t,General		
1357	18259	AI236601	h,v		
1359	17248	Al236635	o,aa		S-phase kinase-associated protein 1A (p19A), transcription elongation factor B (SIII), polypeptide 1 (15 kDa),-like, transcription elongation factor B (SIII), polypeptide 1-like
1360	16859	AI236753	t,General		abrama and Conservation from 4
1361 1362	5208 24388	AI236754 AI236772	g e,General		chromosome 8 open reading frame 1
1363	15850	Al236795	n,v,w		EST, Highly similar to HS9B RAT HEAT SHOCK PROTEIN HSP 90-BETA [R.norvegicus], EST, Weakly similar to HHMS84 heat shock protein 84 - mouse [M.musculus], ESTs, Highly similar to T46243 hypothetical protein DKFZp761K0511.1 [H.sapiens], expressed sequence AL022974, expressed sequence C81438, heat shock 90kD protein 1, beta, heat shock protein, 84 kDa 1
1364	14800	AI236856	w		
1366	11404	Al237002	m		Human DNA sequence from clone RP5-1057D4 on chromosome 20 Contains a spermidine synthase (SPDSY) pseudogene, a CpG island, ESTs, STSs and GSSs, spermidine synthase
4007	40454	A1007040			hepatitis B virus x-interacting protein
1367	18151	AI237212	o,General		(9.6kD)
1368	21653	AI237535	t,General		

Sont Sont	TABLE 8	REMUUHI &	<u>KOMOroen≡ Vi</u>	AMONTATION .		Aiiy, Dockei No. 44921-5089000
	Seallo	ia. National	Consent Acel		പ്രത്യേട്ടിലെട്ടുട്ടി	<u>Doc. No. 1793397.1</u>
Interferon-induced protein - rat R.novegicus EST (R.novegicus) EST R.novegicus EST R.novegicus EST R.novegicus EST R.novegicus EST R.novegicus EST R.novegicus EST R.novegicus EST R.novegicus EST R.novegicus EST R.novegicus RIKEN cDNA 1110036C17 gene, interferon induced transmembrane protein 2 (1-8D) RIKEN cDNA 431348R12 gene, interferon induced transmembrane protein 2 (1-8D) RIKEN cDNA 431348R12 gene, interferon induced transmembrane protein 2 (1-8D) RIKEN CDNA 431348R12 gene, interferon induced transmembrane protein 2 (1-8D) RIKEN CDNA 431348R12 gene, interferon induced transmembrane protein 2 (1-8D) RIKEN CDNA 43137724 F.	No.	ldeniiifer	Ref. Seq. ID No.	Model Gode	Name Care	emski rstavijo zvogelomeki
1370 21893 Al237713 I.k.aa KIAA0101 gene product	1369	11208	A1237586	z		[R.norvegicus], ESTs, Moderately similar to JC1241 beta-interferoninduced protein - rat [R.norvegicus], RIKEN cDNA 1110036C17 gene, RIKEN cDNA 4933438K12 gene, interferon induced transmembrane
1371						
STS, Highly similar to MX11 RAT MX1 INTERTING PROTEIN 1 R. norvegicus), MAX dimerization protein, MAX (INTERTING PROTEIN 1 R. norvegicus), MAX dimerization protein, MAX (intertacting protein 1, Max dimerization protein, MAX (intertacting protein), MAX dimerization protein, MAX (intertacting protein), MAX dimerization protein, MAX (intertacting protein), MAX dimerization protein, MAX (intertacting protein), MAX dimerization protein, MAX (intertacting protein), MAX dimerization protein, MAX (intertacting protein), MAX (int				r		sie ie i gene pieddet
EST, Highly similar to S30385 GSa protein H.sapiens , ESTs, Weakly similar to 1717455 ERG-associated protein EST - mouse M.musculus , HLA-B associated transcript 8, SET domain, bifurcated 1, expressed sequence (7707), hypothetical protein FLJ12873, suppressor of variegation 3-9 (Drosophila) homolog 2; hypothetical protein FLJ12873, suppressor of variegation 3-9 (Drosophila) homolog 2; hypothetical protein FLJ23414 17108	1372	3467	Al237835	General		MAX INTERACTING PROTEIN 1 [R.norvegicus], MAX dimerization protein, MAX-interacting protein 1, Max dimerization protein, Max
1374	1373	25840	AI638972	u		
Maintenance deficient 6 (S. cerevisiae), mini chromosome maintenance deficient 6 (S. cerevisiae), minichromosome maintenance deficient (mis5, S. pombe) 6 deficient (mis5	1374	17108	Al639017	n		protein [H.sapiens], ESTs, Weakly similar to T17453 ERG-associated protein ESET - mouse [M.musculus], HLA-B associated transcript 8, SET domain, bifurcated 1, expressed sequence C77070, hypothetical protein FLJ12879, suppressor of variegation 3-9 (Drosophila) homolog
1376					maintenance deficient 6 (S. cerevisiae), minichromosome maintenance deficient	deficient 6 (S. cerevisiae), minichromosome maintenance
1377 19952 Al639108 q,v					(mis5, S. pombe) 6	deficient (mis5, S. pombe) 6
1379 25907						
1381 18533 Al639231 n hypothetical protein, hypothetical protein FLJ20333 ESTs, Moderately similar to dJ63G5. [H.sapiens], RIKEN cDNA 1700034K16 gene, RIKEN cDNA 170003						
ESTs, Moderately similar to dJ63G5.	1379	25907	Al639167	lo,w		hypothetical protein, hypothetical
1382 18353 Al639233 t,aa decorin 5530600M07 gene, decorin 5530600M07 gene, decorin 5530600M07 gene, decorin 5530600M07 gene, decorin 5530600M07 gene, decorin 5530600M07 gene, decorin 5530600M07 gene, decorin 6530600M07 ge	1381	18533	Al639231	n		ESTs, Moderately similar to dJ63G5.3
1384						1700034K16 gene, RIKEN cDNA
1385 20026 Al639354 g					decorin	5530600M07 gene, decorin
1386 25971						
1388 19152 Al639387 U,General EST, Weakly similar to CAQC RAT CALSEQUESTRIN, CARDIAC MUSCLE ISOFORM PRECURSOR [R.norvegicus], calsequestrin 1, calsequestrin 1 (fast-twitch, skeletal muscle), calsequestrin 2, calsequestrin 2, calsequestrin 2 (cardiac muscle) EST, Moderately similar to A Chain A Mdm2 Bound To The Transactivation Domain Of P53 {SUB 17-125 [H.sapiens], mouse double minute 2, human homolog of; p53-binding protein, transformed mouse 3T3 cell double minute 2 1394 20056 Al639504 a,bb,General)
EST, Weakly similar to CAQC RAT CALSEQUESTRIN, CARDIAC MUSCLE ISOFORM PRECURSOR [R. norvegicus], calsequestrin 1, calsequestrin 1 (fast-twitch, skeletal muscle), calsequestrin 2, calsequestrin 2 (cardiac muscle), calsequestrin 2, calsequestrin 2 (cardiac muscle) EST, Moderately similar to A Chain A Mdm2 Bound To The Transactivation Domain Of P53 {SUB 17-125 [H.sapiens], mouse double minute 2, human homolog of; p53-binding protein, transformed mouse 3T3 cell double minute 2 1392 20082 Al639488 i,m double minute 2 1394 20056 Al639504 a,bb,General						
EST, Moderately similar to A Chain A Mdm2 Bound To The Transactivation Domain Of P53 {SUB 17-125 [H.sapiens], mouse double minute 2, human homolog of; p53-binding protein, transformed mouse 3T3 cell double minute 2 1392 20082 Al639488 i,m double minute 2 1394 20056 Al639504 a,bb,General		19152	Alio3a38/	u,General		CALSEQUESTRIN, CARDIAC MUSCLE ISOFORM PRECURSOR [R.norvegicus], calsequestrin 1,
EST, Moderately similar to A Chain A Mdm2 Bound To The Transactivation Domain Of P53 {SUB 17-125 [H.sapiens], mouse double minute 2, human homolog of; p53-binding protein, transformed mouse 3T3 cell double minute 2 1392 20082 Al639488 i,m double minute 2 1394 20056 Al639504 a,bb,General	1390	18338	AI639422	ly .		
1394 20056 Al639504 a,bb,General				i,m		EST, Moderately similar to A Chain A, Mdm2 Bound To The Transactivation Domain Of P53 (SUB 17-125 [H.sapiens], mouse double minute 2, human homolog of; p53-binding protein, transformed mouse 3T3 cell
	1395	4713	Al639518	q	-	

TABLE	3: HUMAN	HOMOFOGRE	NOTATIONS		Aity. Docket No. 44921-5039WO Doc. No. 1793897.1
Seq. (D) No.		ConBenk Ace./ Ref. Seq. ID No.	Model Code	Homologous Cene Neme	Hanologove Elveter Name
1396 1397	14332 7602	AJ001044 AJ001929	bb lk		EST, Moderately similar to T42215 zonadhesin - mouse [M.musculus], EST, Weakly similar to MUC2_HUMAN MUCIN 2 PRECURSOR [H.sapiens], Homo sapiens SIB 297 intestinal mucin (MUC3) mRNA, partial cds, RIKEN cDNA 3110056H04 gene, RIKEN cDNA 4931407G18 gene, Rattus norvegicus podocalyxin mRNA, complete cds, hepatitis A virus cellular receptor 1, hypothetical protein DKFZp434N185, lymphocyte antigen 64, mucin 1, transmembrane, mucin 2, intestinal/tracheal, zonadhesin
139/	7002	MJ00 1929	, , , , , , , , , , , , , , , , , , ,		EST, Weakly similar to CGHU3B
1398	9867	AJ005424	U		collagen alpha 3(IV) chain precursor, long splice form [H.sapiens], EST, Weakly similar to D40750 proline-rich protein PRB1/2S [H.sapiens], RIKEN cDNA 1190004M21 gene, RIKEN cDNA 2410150I18 gene, RIKEN cDNA 6330577E15 gene, expressed sequence AI551093, mitogenactivated protein kinase 7, murine leukemia viral (bmi-1) oncogene homolog, nemo-like kinase, phospholipid scramblase 1
					ESTs, Weakly similar to CLD7 MOUSE CLAUDIN-7 [M.musculus], ESTs,
1400	16351	AJ011811	General	claudin 7	Weakly similar to CLD7 RAT CLAUDIN 7 [R.norvegicus], claudin 10, claudin 15, claudin 16, claudin 7
1401	20116	AJ011969	I,General	growth differentiation factor 15, prostate differentiation factor	
					ESTs, Moderately similar to BMCP_HUMAN BRAIN MITOCHONDRIAL CARRIER PROTEIN-1 [H.sapiens], ESTs, Weakly similar to M20M_HUMAN MITOCHONDRIAL 2- OXOGLUTARATE/MALATE CARRIER PROTEIN [H.sapiens], ESTs, Weakly similar to brain mitochondrial carrier protein BMCP1 [M.musculus], RIKEN cDNA 1810012H11 gene, expressed sequence AW108044, solute carrier family 25 (mitochondrial carrier; adenine nucleotide translocator), member 10, solute carrier family 25 (mitochondrial carrier; dicarboxylate transporter), member 10, solute carrier family 25 (mitochondrial carrier; ornithine transporter), member 15, solute carrier family 25 (mitochondrial carrier; oxoglutarate carrier), member
1402	17635	AJ223355	v,w		11, uncoupling protein 2, mitochondrial

PCT/US02/16173

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Rof. Seq. D.No. Model Codo Name Homelogous cluster Name ESTs, Highly similar to D3D2 R TRANS-ENCYL-COA ISOMERY MITOCHONDRIAL PRECURSO R. Norvegicus, Home sapiens, to dedecencyl-Coenzyme A delta isomerase (3,2 trans-encyl-Coenzyme A delta isomerase), oldecencyl-Coenzyme A delta isomerase, dodecencyl-Coenzyme A delta isomerase, dodecencyl-Coenzyme A delta isomerase, dodecencyl-Coenzyme A delta isomerase, dodecencyl-Coenzyme A delta isomerase (3,2 trans-encyl-Coenzyme A delta isomerase), dodecencyl-Coenzyme A delta isomerase, dodecencyl-Coenzyme A delta isomerase (3,2 trans-encyl-Coenzyme A delta isomerase), dodecencyl-Coenzyme A delta isomerase, dodecen		APG, Clarifoldina ABC Sinchesia	HOWOTOGATE VI			Atty. Docket No. 4/921-5039W0 Doc. No. 1793397.1
TRANS-ENDYL-COA ISOMERS MITOCHONDRIAL PRECURSO R. Norvegicus , homo sapiens, to dodecenoyl-Coenzyme A delta isomerase (3,2 trans-enoyl-Coenzyme A delta isomerase), clone MGC:3903 MAGE:3630566, mRNA, complete of isomerase), dodecenoyl-Coenzyme A delta isomerase (3,2 trans-enoyl-Coenzyme A delta isomerase), dodecenoyl-Coenzyme A delta isomerase), dodecenoyl-Coenzyme A delta isomerase (3,2 trans-enoyl-Coenzyme A delta isomerase), dodecenoyl-Coenzyme A isomerase, dodecenoyl-Coenzyme A isomerase, dodecenoyl-			ConBank Ace <i>l</i> Ref. Seq. ID No.	Model Gode	Kamologous Cene L Kamo	Homologous Gluster Name
ESTs, Weakly similar to ODP21 DIHYDROLIPOAMIDE ACETYLTRANSFERASE COMPONENT OF PYRUVATE DEHYDROGENASE COMPLEX COMPONENT OF PYRUVATE DEHYDROGENASE COMPLEX [R.norvegicus], Mus musculus, (IMAGE:3586777, mRNA, partial Pyruvate dehydrogenase compl Iipoyl-containing component X; binding protein, RIKEN cDNA 4930529008 gene, dihydrolipoa acetyltransferase (E2 componer pyruvate dehydrogenase comple dihydrolipoamide branched chain transacylase (E2 componer pyruvate dehydrogenase complex dehydrogenase complex maple urine disease), dihydrolipoamide branched chain keto acid dehydrogenase complex maple urine disease), dihydrolipoamide branched chain transacylase E2 transport trans					delta isomerase (3,2 trans- enoyl-Coenyme A isomerase), dodecenoyl- Coenzyme A delta isomerase (3,2 trans-enoyl-	IMAGE:3630566, mRNA, complete cds, dodecenoyl-Coenzyme A delta isomerase (3,2 trans-enoyl-Coenzyme A isomerase), dodecenoyl-Coenzyme A delta isomerase (3,2 trans-enoyl-
DIHYDROLIPOAMIDE ACETYLTRANSFERASE COMPONENT OF PYRUVATE DEHYDROGENASE COMPLEX IRANSFERASE COMPONENT COMPO	1403	18686	D00729	g	Coenzyme A isomerase)	
1405						DIHYDROLIPOAMIDE ACETYLTRANSFERASE COMPONENT OF PYRUVATE DEHYDROGENASE COMPLEX [R.norvegicus], Mus musculus, clone IMAGE:3586777, mRNA, partial cds, Pyruvate dehydrogenase complex, lipoyl-containing component X; E3- binding protein, RIKEN cDNA 1600017E01 gene, RIKEN cDNA 4930529008 gene, dihydrolipoamide S acetyltransferase (E2 component of pyruvate dehydrogenase complex), dihydrolipoamide branched chain transacylase (E2 component of
1405				n,w		branched chain transacylase E2
1406				j		
EST, Moderately similar to DYN						
1409 3015	1407	1613	D14076	x		EST, Moderately similar to DYN2 MOUSE DYNAMIN 2 [M.musculus], ESTs, Moderately similar to DYN2 MOUSE DYNAMIN 2 [M.musculus], KIAA0820 protein, dynamin 2
1410 472 D26111 d,s,bb 1412 16233 D29960 j,I EST, Highly similar to PROTEAS SUBUNIT RC6-1 [R.norvegicus] Highly similar to PSA7_HUMAN PROTEASOME SUBUNIT ALPHTYPE 7 (PROTEASOME SUBUN RC6-1) (PROTEASOME SUBUN XAPC7) [H.sapiens], ESTs, High						
1412 16233 D29960 j,i EST, Highly similar to PROTEAS SUBUNIT RC6-1 [R.norvegicus] Highly similar to PSA7_HUMAN PROTEASOME SUBUNIT ALPH TYPE 7 (PROTEASOME SUBUNI RC6-1) (PROTEASOME SUBUNI RC6-1) (PROTEASOME SUBUNI XAPC7) [H.sapiens], ESTs, High					<u> </u>	
EST, Highly similar to PROTEAS SUBUNIT RC6-1 [R.norvegicus] Highly similar to PSA7_HUMAN PROTEASOME SUBUNIT ALPH TYPE 7 (PROTEASOME SUBUNIT ALPH RC6-1) (PROTEASOME SUBUNIT ALPH XAPC7) [H.sapiens], ESTs, High						
RC6-1 [R.norvegicus], ESTs, We similar to PSA7_HUMAN PROTEASOME SUBUNIT ALPHTYPE 7 (PROTEASOME SUBUNIT ALPHTYPE 7 (PROTEASOME SUBUNIX APC7) [H.sapiens], Human DN sequence from clone RP11-180l chromosome 9 Contains a prote subunit pseudogene, ESTs, STS proteasome (prosome, macropain) subunit, alpha type 7, proteasome (prosome, macropain) subunit, alpha type 7, proteasome (prosome, macropain) subunit, at type 7, proteasome (prosome, macropain)					macropain) subunit, alpha type 7, proteasome (prosome, macropain)	PROTEASOME SUBUNIT ALPHA TYPE 7 (PROTEASOME SUBUNIT RC6-1) (PROTEASOME SUBUNIT XAPC7) [H.sapiens], ESTs, Highly similar to PROTEASOME SUBUNIT RC6-1 [R.norvegicus], ESTs, Weakly similar to PSA7_HUMAN PROTEASOME SUBUNIT ALPHA TYPE 7 (PROTEASOME SUBUNIT RC6-1) (PROTEASOME SUBUNIT XAPC7) [H.sapiens], Human DNA sequence from clone RP11-18014 on chromosome 9 Contains a proteasome subunit pseudogene, ESTs, STSs, GSSs and a CpG island, RIKEN cDNA 2410072D24 gene, proteasome (prosome, macropain) subunit, alpha type 7, proteasome (prosome,
1413 9029 D30804 n subunit, alpha type, 7 macropain) subunit, alpha type, 7	1413	9029	D30804	n	subunit, alpha type, 7	macropain) subunit, alpha type, 7

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					<u>Dos. No. 1793897.1</u>
Seq. ID No:	ldentifier	ConEank Ace.// Ref. Seq. ID No.	Model Code	Hamologous Gene Neme	Homologous Cluster Name
					PROTEIN-TYROSINE PHOSPHATASE-LIKE N PRECURSOR [H.sapiens], protein tyrosine phosphatase, receptor type,
1414	1485	D38222	y,z		N, protein tyrosine phosphatase, receptor-type, N
1415	9135	D45247	S	proteasome (prosome, macropain) subunit, beta type 5, proteasome (prosome, macropain) subunit, beta type, 5	EST, Moderately similar to PRCE RAT PROTEASOME EPSILON CHAIN PRECURSOR [R.norvegicus], RIKEN cDNA 5830406J20 gene, proteasome (prosome, macropain) subunit, beta type 5, proteasome (prosome, macropain) subunit, beta type, 5
1416	16354	D50564	u		proteasome (prosome, macropain)
1417	1884_	D50695	l,m,bb		26S subunit, ATPase, 4
1418	21147	D63772	General	solute carrier family 1 (neuronal/epithelial high affinity glutamate transporter, system Xag), member 1, solute carrier family 1, member 1	
					CDP-diacylglycerolinositol 3- phosphatidyltransferase
1419 1420	826 25306	D82928 D84485	f u		(phosphatidylinositol synthase)
1421	18867 22543	D88250 H31117	t r,v,General		EST, Moderately similar to MAS2_HUMAN MANNAN-BINDING LECTIN SERINE PROTEASE 2 PRECURSOR [H.sapiens], EST, Weakly similar to JC6554 probable serine proteinase [R.norvegicus], ESTs, Weakly similar to JC6554 probable serine proteinase [R.norvegicus], MASP-2 protein, complement component 1, r subcomponent, complement component 1, r subcomponent 1, s subcomponent, mannan-binding lectin serine protease 1, mannan-binding lectin serine protease 2
1424 1425	12360 20514	H31456 H31489	h,j		
1426	11358	H31610			DNA segment, Chr 16, Indiana University Medical 21, expressed, ESTs, Highly similar to JC5020 tetratricopeptide repeat protein [H.sapiens], ESTs, Moderately similar to JW0059 mtprd protein - mouse [M.musculus], PRO1880 protein, g1- related zinc finger protein, tetratricopeptide repeat domain,
1426	4360	H31813	h bb,General		tetratricopeptide repeat domain 3 DKFZP586B1621 protein
1428	9343	H32169	1		EST, Moderately similar to COF1_HUMAN COFILIN, NON- MUSCLE ISOFOR [H.sapiens], cofilin 1 (non-muscle), cofilin 1, non-muscle, cofilin 2 (muscle), cofilin 2, muscle
1429	4386	H33093	h,w		Z (massio), comm z, massie
1430	4415	H33636	h		EST- Weekly similar to 1500 Million
1431	15374	H34186	I		ESTs, Weakly similar to IF39_HUMAN EUKARYOTIC TRANSLATION INITIATION FACTOR 3 SUBUNIT 9 [H.sapiens]

TABLE 3	HUMAN.	HOMOFOGAE VIX	RAGITATIONS		Atty: Docket No. 44921-5089W0
		locatorati can i			<u> </u>
Seq. ID		Cenbank Ace./ Ref. Seq. ID No.		Homologous Gene News	Homologous Gluster Name
1432	17159	J00797	u.General	Keme	ESTs, Highly similar to A23035 tubulin alpha chain [H.sapiens], tubulin alpha 1, tubulin alpha 2, tubulin alpha 3, tubulin alpha 6, tubulin, alpha 2, tubulin, alpha 3, tubulin, alpha 3, tubulin, alpha, lubiquitous
1433	16260	J01878	f		
			LL	branched chain keto acid dehydrogenase E1, alpha polypeptide (maple syrup urine disease), branched chain ketoacid dehydrogenase E1, alpha	
1434 1435	17284 15017	J02827 J03752	bb n	polypeptide	
1436	44	J03819	p.s	thyroid hormone receptor beta, thyroid hormone receptor, beta (avian erythroblastic leukemia viral (v-erb-a) oncogene homolog 2)	
				glutathione S-transferase M2 (muscle), glutathione S-	
1437	21014	J03914	e,r,General	transferase, mu 2	transferase, mu 1
1438	20429	J05035	f	steroid 5 alpha-reductase 1, steroid-5-alpha- reductase, alpha polypeptide 1 (3-oxo-5 alpha-steroid delta 4- dehydrogenase alpha 1)	steroid-5-alpha-reductase, alpha polypeptide 1 (3-oxo-5 alpha-steroid delta 4-dehydrogenase alpha 1)
	10.17	10.5101		glutamate-cysteine ligase,	glutamate-cysteine ligase, catalytic
	10464 20149	J05510 K03243	n,u,General	inositol 1,4,5-triphosphate receptor 1, inositol 1,4,5-triphosphate receptor, type 1	subunit ESTs, Highly similar to A55713 inositol 1,4,5-triphosphate receptor type 1 [H.sapiens], ESTs, Weakly similar to IP3R MOUSE INOSITOL 1,4,5- TRISPHOSPHATE-BINDING PROTEIN TYPE 1 RECEPTOR [M.musculus], expressed sequence Al528790, inositol 1,4,5-triphosphate receptor 1, inositol 1,4,5-triphosphate receptor, type 1
1441	20149	KU3243	q		
		K03249	q		ESTs, Highly similar to ECHP_HUMAN PEROXISOMAL BIFUNCTIONAL ENZYME [H.sapiens], enoyl-Coenzyme A, hydratase/3-hydroxyacyl Coenzyme A dehydrogenase
		L00124	w		
	2048	L00382	k,x	· · · · · · · · · · · · · · · · · · ·	
		L04619 L14002	S		
		L14002	p t		
		L14004	c,p		
			General	phosphoinositide-specific phospholipase C-beta 1, phospholipase C, beta 1	ESTs, Highly similar to KIAA0581 protein [H.sapiens]
		L14323	General		
		L14937	у	01.01	01.01
1451	25369		k	2',3'-cyclic nucleotide 3' phosphodiesterase, cyclic nucleotide phosphodiesterase 1 phosphofructokinase,	2',3'-cyclic nucleotide 3' phosphodiesterase, cyclic nucleotide phosphodiesterase 1, hypothetical gene CG018

TABLE	38 HUMAN	HONOLOGUEA	SKIÐITATIÐKE		Atty, Docket No. 44921-5039W0 Doc. No. 1793597.1
Seg. (D) No.		ConBenk Ace./ Ref. Seq. ID No.		Homologous Cene Name	Homologous Cluster Name
1453	12058	L25387	h	phosphofructokinase, platelet	ESTs, Highly similar to JC2055 6-phosphofructokinase [H.sapiens], ESTs, Weakly similar to JC2055 6-phosphofructokinase [H.sapiens], Musmusculus adult male stomach cDNA, RIKEN full-length enriched library, clone:2210403E17, full insert sequence, expressed sequence AA407869, phosphofructokinase, liver, B-type, phosphofructokinase, muscle, phosphofructokinase, platelet
1455	21146	L35558	General	solute carrier family 1 (neuronal/epithelial high affinity glutamate transporter, system Xag), member 1, solute carrier family 1, member 1	
1456	106	L37203	w	, , , , , , , , , , , , , , , , , , ,	
1458	13682	L38482	f,j,k,m,z	-	
1459	6405	L38615	p	glutathione synthetase	
1461	15189	M11794	n,v		
1462	17086	M13011	j		
1464	21053	M15481	0		
1465	25405	M18330	j,l		
1466 1468	25415 14967	M19648 M22366	a w		
1469	20481	M22631	bb		
1471	15048	M24542	q		REDUCTASE IRON-SULFUR SUBUNIT, MITOCHONDRIAL PRECURSOR [R.norvegicus], EST, Weakly similar to UCRI_HUMAN UBIQUINOL-CYTOCHROME C REDUCTASE IRON-SULFUR SUBUNIT, MITOCHONDRIAL PRECURSOR [H.sapiens], ESTs, Moderately similar to UCRI_HUMAN UBIQUINOL-CYTOCHROME C REDUCTASE IRON-SULFUR SUBUNIT, MITOCHONDRIAL PRECURSOR [H.sapiens], Human DNA sequence from clone RP1-228J4 on chromosome 6 Contains a pseudogene similar to UQCRFS1 (ubiquinol-cytochrome c reductase, Rieske iron-sulfur polypeptide 1), ESTs, an STS and GSSs, RIKEN cDNA 4430402G14 gene, expressed sequence Al875505, ubiquinol- cytochrome c reductase, Rieske iron- sulfur polypeptide 1
				cytochrome P450, subfamily IV B, polypeptide 1, cytochrome P450, subfamily IVB, polypeptide	Mus musculus, Similar to cytochrome P450, 4a10, clone MGC:25972 IMAGE:4240359, mRNA, complete cds, RIKEN cDNA A230105L22 gene, cytochrome P450, 4a10, cytochrome P450, 4a14, cytochrome P450,
1472	20921	M29853	m	1	subfamily IVA, polypeptide 11 cytochrome P450, subfamily IIA
1473	1224	M31931	ļ.,		(phenobarbital-inducible), polypeptide 13
1474	15579	M33648	q		10
	1	M33648	17		

TABLE 8	HUMAN	HOMOFOGAE W	EXIONATION		Atty, Docket No. 44921-5039W0
Seg. ID No.	(demilier	ConBink Acel Rol Son ID No.	Modal (Code)	Homologous Gene Name	Doc. No. 1793397.1 Homologous Cluster Name
	Chamana de la caracterí				EST, Moderately similar to 60S RIBOSOMAL PROTEIN L35 [R.norvegicus], EST, Moderately similar to G01477 ribosomal protein L35 [H.sapiens], EST, Weakly similar to 60S RIBOSOMAL PROTEIN L35 [R.norvegicus], ESTs, Highly similar to G01477 ribosomal protein L35 [H.sapiens], Human DNA sequence from clone RP1-34P24 on chromosome 22 Contains a pseudogene similar to ribosomal protein L35, ESTs, STSs and GSSs, RIKEN cDNA 2410039E09 gene,
1475	17211	M34331	g,n,q,v		ribosomal protein L35 Homo sapiens clone HQ0582,
1476	20699	M35601	b,x,bb	fibrinogen, A alpha polypeptide	expressed sequence Al303526, fibrinogen, A alpha polypeptide, fibrinogen, gamma polypeptide Homo sapiens clone HQ0582, expressed sequence Al303526,
1476	20700	M35601	b,t,bb	fibrinogen, A alpha polypeptide	fibrinogen, A alpha polypeptide, fibrinogen, gamma polypeptide H.sapiens DMA, DMB, HLA-Z1, IPP2,
	0000	1100454			LMP2, TAP1, LMP7, TAP2, DOB, DQB2 and RING8, 9, 13 and 14
1477 1479	9223 1585	M36151 M57728	j,m,y	-	genes, expressed sequence Al845868
4490	24044	M58040			DKFZp434M2227 (from clone DKFZp434M2227), Homo sapiens prostate-specific membrane antigen PSM mRNA, exon 6 alternative splice variant, partial cds, RIKEN cDNA 2610028K12 gene, folate hydrolase, transferrin receptor, transferrin receptor (p90, CD71), transferrin receptor 2
1480	24844 25057	M58495	h		receptor 2
1482	457	M60666	d,General	tropomyosin 1 (alpha), tropomyosin 1, alpha	ESTs, Moderately similar to alphatropomyosin slow [M.musculus], tropomyosin 4
1484	5733 4198	M81855	i,k,aa m	ATP-binding cassette, sub- family B (MDR/TAP), member 1, ATP-binding cassette, sub-family B (MDR/TAP), member 1B	[R.norvegicus], ESTs, Weakly similar to MDR1 MOUSE MULTIDRUG RESISTANCE PROTEIN 1 [M.musculus] ESTs, Highly similar to A41734 betagalactoside alpha-2,6-sialyltransferase [H.sapiens] ESTs, Highly similar to A41734 betagalactoside alpha-2,6-sialyltransferase
1485	4199 24651	M83143 M83678	m k,x,z		[H.sapiens] RAB13, member RAS oncogene family, RIKEN cDNA 0610007N03 gene, expressed sequence AW107754

TABLES	MAMUEL	HOWOFOGME W	SKIONTATIONI		Aiiy. Docket No. 44921-5039WO Doc. No. 1793397.1
809. [D No. ' '	ldentifier	ConBank Acc./ Ref. Seq. ID No.	Model Code	Homologous Gene Neme	Homologous Cluster Name
1487	1430	M84648	General	dopa decarboxylase, dopa decarboxylase (aromatic L- amino acid decarboxylase)	EST, Highly similar to AROMATIC-L-AMINO-ACID DECARBOXYLASE [R.norvegicus], EST, Moderately similar to AROMATIC-L-AMINO-ACID DECARBOXYLASE [R.norvegicus], Mus musculus 10 days embryo cDNA, RIKEN full-length enriched library, clone:2610109O21, full insert sequence, dopa decarboxylase, dopa decarboxylase (aromatic L-amino acid decarboxylase)
1488	25467	M93297	C	animo acid decarboxylase)	(decarboxylase)
					ESTs, Highly similar to NTG3 MOUSE SODIUM- AND CHLORIDE-DEPENDENT GABA TRANSPORTER 3 [M.musculus], ESTs, Weakly similar to NTG3_HUMAN SODIUM- AND CHLORIDE-DEPENDENT GABA TRANSPORTER 3 [H.sapiens], expressed sequence AA589632, solute carrier family 6 (neurotransmitter transporter, GABA),
1489	729	M95762	a,y		member 13
1490	23698	NM 012489	g.		DNA segment, Chr 9, ERATO Doi 25, expressed, Homo sapiens clone 23623 mRNA, partial cds, Homo sapiens, Similar to Acetyl-CoA acyltransferase, 3-oxo acyl-CoA thiolase A, peroxisomal, clone MGC:18173 IMAGE:4155289, mRNA, complete cds, Homo sapiens, Similar to acetyl-Coenzyme A acyltransferase 1 (peroxisomal 3-oxoacyl-Coenzyme A thiolase), clone MGC:23127 IMAGE:4908159, mRNA, complete cds, RIKEN cDNA 0610011L04 gene, acetyl-Coenzyme A acyltransferase 1 (peroxisomal 3-oxoacyl-Coenzyme A thiolase), expressed sequence Al255831, expressed sequence Al265397, t-complex protein 1, related sequence 1
1490	23698	NM_012489	q		DNA segment, Chr 9, ERATO Doi 25, expressed, Homo sapiens clone 23623
1490	23699	NM_012489			mRNA, partial cds, Homo sapiens, Similar to Acetyl-CoA acyltransferase, 3-oxo acyl-CoA thiolase A, peroxisomal, clone MGC:18173 IMAGE:4155289, mRNA, complete cds, Homo sapiens, Similar to acetyl-Coenzyme A acyltransferase 1 (peroxisomal 3-oxoacyl-Coenzyme A thiolase), clone MGC:23127 IMAGE:4908159, mRNA, complete cds, RIKEN cDNA 0610011L04 gene, acetyl-Coenzyme A acyltransferase 1 (peroxisomal 3-oxoacyl-Coenzyme A thiolase), expressed sequence Al255831, expressed sequence Al265397, t-complex protein 1, related sequence 1

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Seq. ID No.	ldentifier	ConBank Ace./ Roi. Seq. ID No.	Model Gode	Homologous Cana Name	Homologous Gluster Name
and the second second				aldolase 1, A isoform, aldolase A, fructose-	EST, Moderately similar to ADHUA fructose-bisphosphate aldolase [H.sapiens], EST, Weakly similar to ADHUA fructose-bisphosphate aldolase [H.sapiens], EST, Weakly similar to 139435 fructose-bisphosphate aldolase [H.sapiens], ESTs, Moderately similar to aldolase A [M.musculus], Homo sapiens, aldolase A [M.musculus], Homo sapiens, aldolase 1, A isoform, clone MGC:18171 IMAGE:4155253, mRNA, complete cds, RIKEN cDNA 4933425L11 gene, aldolase 1, A isoform, aldolase 3, C isoform, aldolase A, fructose-
1491	7062	NM_012495	q	bisphosphate	bisphosphate
1492	15511	NM 012498	lu	aldo-keto reductase family 1, member B1 (aldose reductase), aldo-keto reductase family 1, member B3 (aldose reductase)	
1494	7427	NM_012515	General	benzodiazapine receptor (peripheral), benzodiazepine receptor, peripheral cholinergic receptor, muscarinic 3, cholinergic receptor, muscarinic 3,	Human DNA sequence from clone 34B21 on chromosome 6p12.1-21.1. Contains part of a gene for a novel protein with ZU5 domain similar to part of Tight Junction Protein ZO1 (TJP1) and UNC5 Homologs, the gene for a novel BZRP (peripheral benzodiazapine receptor (MBR,PBR, PBKS, IBP, Isoquinoline-binding protein)) LIKE protein, the gene for a novel protein similar to part of APOBEC1 (Phorbolin 1, Apolipoprotein B mRNA editing protein), and the NFYA gene for nuclear transcription factor Y, alpha (CCAAT-Binding transcription factor subunit B, CBF-B, CAAT-Box DNA binding protein subunit A). Contains ESTs, STSs, GSSs, two putative CpG islands and a ca repeat polymorphism, RIKEN cDNA 2510027D20 gene, benzodiazapine receptor (peripheral), benzodiazapine receptor, peripheral cholinergic receptor, muscarinic 1, CNS, cholinergic receptor, muscarinic 3, cardiac
1495	24433	NM_012527	i	cardiac	cardiac
1496	4467	NM_012529	d	creatine kinase, brain	EST, Moderately similar to CREATINE KINASE, B CHAIN [R.norvegicus], EST, Weakly similar to KIHUCB creatine kinase [H.sapiens], creatine kinase, brain
1497	16520	NM 012532	General	ceruloplasmin,	DNA segment, Chr 3, ERATO Doi 555, expressed, EST, Highly similar to 1012298A factor VIIIC [H.sapiens], ESTs, Weakly similar to CERU MOUSE CERULOPLASMIN PRECURSOR [M.musculus], ESTs, Weakly similar to CERU RAT CERULOPLASMIN PRECURSOR [R.norvegicus], ESTs, Weakly similar to KUHU ferroxidase [H.sapiens], Hermansky-Pudlak syndrome 3, ceruloplasmin, ceruloplasmin

TABLES	: HUMAN	IHOMOITOGNE W			Anny, Docket No. 44921-5139Wo Doc. No. 1793597.1
Seq. 10 : No:	ldentifier	Centent Acel Ref. Seq. ID No.	Model Code	Homologous Gene Neme	Homologous Gluster Name
1498	225	NM 012544	x,z	angiotensin I converting enzyme (peptidyl- dipeptidase A) 1, angiotensin converting enzyme	ESTs, Highly similar to A31759 peptidyl-dipeptidase A [H.sapiens], ESTs, Highly similar to JC2038 peptidyl-dipeptidase A [R.norvegicus], Mus musculus adult male testis cDNA, RIKEN full-length enriched library, clone:4933424D04, full insert sequence, RIKEN cDNA 2010305L05 gene, angiotensin I converting enzyme (peptidyl-dipeptidase A) 1, angiotensin converting enzyme
	·			dopa decarboxylase, dopa decarboxylase (aromatic L-	EST, Highly similar to AROMATIC-L-AMINO-ACID DECARBOXYLASE [R.norvegicus], EST, Moderately similar to AROMATIC-L-AMINO-ACID DECARBOXYLASE [R.norvegicus], Mus musculus 10 days embryo cDNA, RIKEN full-length enriched library, clone:2610109021, full insert sequence, dopa decarboxylase, dopa decarboxylase (aromatic L-amino acid
1499	1431	NM_012545	General	amino acid decarboxylase)	decarboxylase) ESTs, Weakly similar to I53869 zinc finger protein - mouse [M.musculus],
1500	23868	NM_012551	I,m,v,General	early growth response 1	early growth response 1, repressor of GATA, testis zinc finger protein
1500	23872	NM_012551	I,v,cc,General	early growth response 1	ESTs, Weakly similar to I53869 zinc finger protein - mouse [M.musculus], early growth response 1, repressor of GATA, testis zinc finger protein ESTs, Weakly similar to I53869 zinc finger protein - mouse [M.musculus],
1500	23869	NM_012551	v,General	early growth response 1	early growth response 1, repressor of GATA, testis zinc finger protein
1501	19407	NM_012554	z	enolase 1, (alpha), enolase 1, alpha non-neuron	EST, Moderately similar to ALPHA ENOLASE [R.norvegicus], EST, Weakly similar to A29170 phosphopyruvate hydratase [H.sapiens], Homo sapiens cDNA FLJ12774 fis, clone NT2RP2001663, moderately similar to ENOLASE (EC 4.2.1.11), enolase 1, alpha, enolase 1, alpha non-neuron, enolase alpha, lung-specific, expressed sequence Al427012
				enolase 1, (alpha), enolase	EST, Moderately similar to ALPHA ENOLASE [R.norvegicus], EST, Weakly similar to A29170 phosphopyruvate hydratase [H.sapiens], Homo sapiens cDNA FLJ12774 fis, clone NT2RP2001663, moderately similar to ENOLASE (EC 4.2.1.11), enolase 1, alpha, enolase 1, alpha non-neuron, enolase alpha, lung-specific, expressed sequence
1501	19408	NM_012554	n,s,y,z	1, alpha non-neuron	Al427012 E26 avian leukemia oncogene 1, 5'
1502	21836	NM_012555	k	E26 avian leukemia oncogene 1, 5' domain, v- ets avian erythroblastosis virus E26 oncogene homolog 1	domain, ESTs, Moderately similar to FLI1 MOUSE RETROVIRAL INTEGRATION SITE PROTEIN FLI-1 [M.musculus], FEV protein, Rattus norvegicus ETS domain transcription factor Pet-1 mRNA, complete cds, vets avian erythroblastosis virus E26 oncogene homolog 1

TABLES	e <mark>ihium</mark> an	HOMOLOGUE AN	(EXOTATOXE		Atty. Docket No. 44921-5039W6 Doc. No. 1793897.1
539, ID No. 🏥	ldenijier	ConBank Acel. Ref. Seq. ID No.		Homologous Care Name	Honologous Glister Name
1503 1504	16895 25317	NM_012558 NM_012559	g,s bb		EST, Weakly similar to FRUCTOSE- 1,6-BISPHOSPHATASE ISOZYME 2 [M.musculus], fructose bisphosphatase 1, fructose bisphosphatase 2, fructose- 1,6-bisphosphatase 1, fructose-1,6- bisphosphatase 2
1504	25317	NM_012559	DD		EST, Moderately similar to FGHUGB
1504	6477	NM 012559	b,bb	fibrinogen, gamma polypeptide	fibrinogen gamma-B chain precursor [H.sapiens], EST, Moderately similar to Recombinant Human Gamma-Fibrinogen Carboxyl Terminal Fragment [H.sapiens], ESTs, Weakly similar to FIBG RAT FIBRINOGEN GAMMA-A-B CHAIN PRECURSOR [R.norvegicus], RIKEN cDNA 1110007F23 gene, expressed sequence Al303526, fibrinogen, gamma polypeptide, fibrinogen-like 1, ficolin A, ficolin B
				fibrinogen, gamma	EST, Moderately similar to FGHUGB fibrinogen gamma-B chain precursor [H.sapiens], EST, Moderately similar to Recombinant Human Gamma-Fibrinogen Carboxyl Terminal Fragment [H.sapiens], ESTs, Weakly similar to FIBG RAT FIBRINOGEN GAMMA-A/-B CHAIN PRECURSOR [R.norvegicus], RIKEN cDNA 1110007F23 gene, expressed sequence Al303526, fibrinogen, gamma polypeptide, fibrinogen-like 1,
1504	6478	NM_012559	bb	polypeptide	ficolin A, ficolin B follistatin, follistatin-like 3, follistatin- like 3 (secreted glycoprotein),
1505	11731	NM_012561	k	follistatin	transmembrane protein with EGF-like and two follistatin-like domains 1
1000	11731	1410_012301	<u> </u>	group specific component,	and two lonistatii-like domains i
				group-specific component	
1507	4254	NM_012564	а	(vitamin D binding protein) H1 histone family, member	
1508	16026	NM_012578	r	10	H1 histone family, member 0
	•			H1 histone family, member	· · · · · · · · · · · · · · · · · · ·
1508	16024	NM_012578	r	0	H1 histone family, member 0
1508	16025	NM_012578	r	H1 histone family, member 0	H1 histone family, member 0
				heme oxygenase	
1509	16080	NM_012580	g,m	(decycling) 1	heme oxygenase (decycling) 1
1510	15098	NM_012588	bb	insulin-like growth factor binding protein 3	insulin-like growth factor binding protein 3, protease, serine, 11 (Igf binding)
1511	4450	NM_012592	bb	isovaleryl Coenzyme A dehydrogenase, isovaleryl coenzyme A dehydrogenase	RIKEN cDNA 1300003O09 gene, RIKEN cDNA 2310016C19 gene, acetyl-Coenzyme A dehydrogenase, short chain, isovaleryl Coenzyme A dehydrogenase, isovaleryl coenzyme A dehydrogenase RIKEN cDNA 1300003O09 gene,
1511	4451	NM_012592	i,bb	isovaleryl Coenzyme A dehydrogenase, isovaleryl coenzyme A dehydrogenase	RIKEN CDNA 1300003009 gene, RIKEN cDNA 2310016C19 gene, acetyl-Coenzyme A dehydrogenase, short chain, isovaleryl Coenzyme A dehydrogenase, isovaleryl coenzyme A dehydrogenase

TABLE	3: HUMAN	HOMOLOGUE AN	RIOTRATIONS		Airy. Docket No. 44921-5039000 Doc. No. 1793397.
Sog. (D No.	ldentifter	Consonk Aced Ref. Seq. ID No.		Homologous Gene Name	Homologous Cluster Name
	1.0 Indianate Management Street Management	Decision of the Conference o		isovaleryl Coenzyme A dehydrogenase, isovaleryl	RIKEN cDNA 130003009 gene, RIKEN cDNA 2310016C19 gene, acetyl-Coenzyme A dehydrogenase, short chain, isovaleryl Coenzyme A
1511	4452	NM_012592	bb	coenzyme A dehydrogenase	dehydrogenase, isovaleryl coenzyme A dehydrogenase
1512	17198	NM_012593	a,x	kallikrein 1, renal/pancreas/salivary, kallikrein 6	kallikrein 1, renal/pancreas/salivary, kallikrein 21, kallikrein 24, kallikrein 27 kallikrein 5
1512	17197	NM_012593	x	kallikrein 1, renal/pancreas/salivary, kallikrein 6	kallikrein 1, renal/pancreas/salivary, kallikrein 21, kallikrein 24, kallikrein 27 kallikrein 5
1513	18749	NM_012600	a,h	malic enzyme 1, NADP(+)- dependent, cytosolic, malic enzyme, supernatant	
1514	2628	NM_012603	General	myelocytomatosis oncogene, v-myc avian myelocytomatosis viral oncogene homolog myelocytomatosis oncogene, v-myc avian	RIKEN cDNA 2900002K07 gene, myelocytomatosis oncogene, v-myc
1514	2629	NM 012603	x,General	myelocytomatosis viral oncogene homolog	avian myelocytomatosis viral oncogene homolog
1515	16849	NM_012608	n,o,q	membrane metallo endopeptidase, membrane metallo-endopeptidase (neutral endopeptidase, enkephalinase, CALLA, CD10)	endothelin converting enzyme-like 1, expressed sequence AW322500, mel transforming oncogene-like 1, membrane metallo endopeptidase, membrane metallo-endopeptidase (neutral endopeptidase, enkephalinase, CALLA, CD10), membrane metallo-endopeptidase-like 2
1517		NM_012620	General	serine (or cysteine) proteinase inhibitor, clade E (nexin, plasminogen activator inhibitor type 1), member 1	serine (or cysteine) proteinase inhibitor, clade E (nexin, plasminogen activator inhibitor type 1), member 1
1518	24568	NM_012630	General	prolactin receptor	cytokine receptor-like factor 1, interleukin 13 receptor, alpha 2, interleukin 5 receptor, alpha, prolactin receptor, prolactin receptor related sequence 1
1518	24566	NM_012630	General	prolactin receptor	cytokine receptor-like factor 1, interleukin 13 receptor, alpha 2, interleukin 5 receptor, alpha, prolactin receptor, prolactin receptor related sequence 1
1519	18553	NM_012631	k	prion protein, prion protein (p27-30) (Creutzfeld-Jakob disease, Gerstmann- Strausler-Scheinker syndrome, fatal familial insomnia)	prion protein, prion protein (p27-30) (Creutzfeld-Jakob disease, Gerstmann Strausler-Scheinker syndrome, fatal familial insomnia)
1010	10000	1101_012001	N	ingollilla)	ESTs, Highly similar to TPHUN1 protein-tyrosine-phosphatase
1520	1844	NM_012637	General		[H.sapiens], protein tyrosine phosphatase, non-receptor type 1 expressed sequence D19352, renin,
1521 1522 1523	24668 18632 25435	NM_012642 NM_012645 NM_012647	f a g	renin, renin 1 structural	renin 1 structural, renin 2 tandem duplication of Ren1
1524	9423	NM_012649	b,cc	syndecan 4, syndecan 4 (amphiglycan, ryudocan)	syndecan 4, syndecan 4 (amphiglycan ryudocan)

TABLES	HUMAN.	HOMOLOGUEAN	INOTATIONS		Ality. Docket No. 44921-5039W0 Doc. No. 1793897.1
Seg. (D) No. : '	ldentifier	GenBank Ace./ Ref. Seq. 10 No.	Model Gode	Honologous Cons Neme	Homologous Cluster Name
					ESTs, Weakly similar to NAH1 MOUSE SODIUM/HYDROGEN EXCHANGER 1 [M.musculus], Homo sapiens mRNA; cDNA DKFZp434D0818 (from clone DKFZp434D0818), RIKEN cDNA 0610040A22 gene, RIKEN cDNA 6430709P13 gene, expressed
1525	24496	NM 012654	n	solute carrier family 9 (sodium/hydrogen exchanger), isoform 3, solute carrier family 9 (sodium/hydrogen exchanger), member 3	sequence AI182282, solute carrier family 9 (sodium/hydrogen exchanger), isoform 3, solute carrier family 9 (sodium/hydrogen exchanger), isoform 6, solute carrier family 9 (sodium/hydrogen exchanger), member 1
				overlangery, member c	DNA segment, Chr 14, University of California at Los Angeles 3, EST, Moderately similar to A41386 clusterin precursor [H.sapiens], ESTs, Moderately similar to A41386 clusterin precursor [H.sapiens], clusterin, clusterin (complement lysis inhibitor, SP-40,40, sulfated glycoprotein 2, testosterone-repressed prostate message 2, apolipoprotein J),
1526	7101	NM_012679	x,bb,General		expressed sequence AI893575 cytochrome P450, 2a12, cytochrome P450, subfamily IIA (phenobarbital-inducible), polypeptide 13, cytochrome P450, subfamily IIA (phenobarbital-inducible), polypeptide 7, pseudogene 1, cytochrome P450, subfamily IIA (phenobarbital-inducible), polypeptide
1527	24707	NM_012693			7, pseudogene 2 EST, Weakly similar to KNG_MOUSE KININOGEN PRECURSOR
1528	1850	NM_012696	t	kininogen	[CONTAINS: BRADYKININ] [M.musculus], kininogen EST, Weakly similar to KNG_MOUSE KININOGEN PRECURSOR
1528	1854	NM_012696	t	kininogen	[CONTAINS: BRADYKININ] [M.musculus], kininogen EST, Weakly similar to OCN2 MOUSE ORGANIC CATION/CARNITINE TRANSPORTER 2 [M.musculus], ESTs, Highly similar to OCN2_HUMAN ORGANIC CATION/CARNITINE TRANSPORTER 2 [H.sapiens], solute carrier family 22 (organic cation transporter), member 1, solute carrier family 22 (organic cation transporter), member 4, solute carrier family 22 (organic cation transporter), member
1529	1603	NM_012697	General		5, solute carrier family 22 (organic cation transporter), member 9 ESTs, Highly similar to HXK1 RAT
1530	1372	NM_012734	u	hexokinase 1	HEXOKINASE, TYPE I [R.norvegicus], hexokinase 1, hypothetical protein FLJ22761
1531	1478	NM_012744	bb,General	TIONORIHAGO I	expressed sequence C79630, methylcrotonoyl-Coenzyme A carboxylase 1 (alpha), pyruvate carboxylase, pyruvate decarboxylase

TABLE	S HUMAN	HOMOFOGUEAN			Atty: Docket No. 44921-508900 Doc. No. 1798897.1
8eg. (D. No.	ldentifier	Conbank Acc./ Ref. Seq. ID No.	Model Code	Homologous Gere Name	Homologous Cluster Name
1532 1533	343 8829	NM_012747 NM_012749	h,t General		Signal transducer and activator of transcription 1, signal transducer and activator of transcription 1, signal transducer and activator of transcription 1, 91kD, signal transducer and activator of transcription 3, signal transducer and activator of transcription 3 (acute-phase response factor)
	0020		00,10,0	CD24 antigen (small cell	CD24 antigen (small cell lung
1534	20828	NM_012752	General	lung carcinoma cluster 4 antigen), CD24a antigen	carcinoma cluster 4 antigen), CD24a antigen
1534	20829	NM 012752	i,General	CD24 antigen (small cell lung carcinoma cluster 4 antigen), CD24a antigen	CD24 antigen (small cell lung carcinoma cluster 4 antigen), CD24a antigen
				CD24 antigen (small cell lung carcinoma cluster 4	CD24 antigen (small cell lung carcinoma cluster 4 antigen), CD24a
1534	20830	NM_012752	i,General	antigen), CD24a antigen	antigen ESTs, Highly similar to 1312358A IGF
1535	15174	NM_012756	b		Il receptor [H.sapiens], expressed sequence Al661837, insulin-like growth factor 2 receptor EST, Weakly similar to ZF37 RAT
1536	21685	NM 012760	j,m,n		ZINC FINGER PROTEIN 37 (ZFP-37) [R.norvegicus], ESTs, Weakly similar to S59069 Z13 protein - mouse [M.musculus], Homo sapiens chromosome 19, BAC 273239 (CIT-B-320G13), Homo sapiens, clone MGC:23189 IMAGE:4854518, mRNA, complete cds, RIKEN cDNA 2410081M15 gene, RIKEN cDNA 2610019F01 gene, RIKEN cDNA 2810011C24 gene, hypothetical protein FLJ12488, hypothetical zinc finger protein MGC2396, zinc finger protein 37, zinc finger protein homologous to Zfp37 in mouse
					CARD only protein, ESTs, Moderately similar to A56084 interleukin-1beta converting enzyme beta isozyme [H.sapiens], ESTs, Weakly similar to interleukin-1beta converting enzyme gamma isozyme [H.sapiens], Homo sapiens mRNA; cDNA DKFZp586A181 (from clone DKFZp586A181); partial cds, ICEBERG caspase-1 inhibitor, caspase 1, caspase 1, apoptosisrelated cysteine protease (interleukin
1537	18068	NM_012762	t		1, beta, convertase) ESTs, Weakly similar to ANPA MOUSE ATRIAL NATRIURETIC PEPTIDE RECEPTOR A PRECURSOR [M.musculus], guanylate cyclase 1, soluble, beta 2, guanylate cyclase 1, soluble, beta 3,
1538 1539	1246	NM_012770 NM_012776	a,General	adrenergic receptor kinase, beta 1, adrenergic, beta, receptor kinase 1	Guardiale Cyclase 1, Soluble, Deta 3, natriuretic peptide receptor 1 G protein-coupled receptor kinase 2, groucho gene related (Drosophila), G protein-coupled receptor kinase 5, G protein-coupled receptor kinase 6, adrenergic, beta, receptor kinase 1, rhodopsin kinase

TABLES	HUMAN	HOMOLOGUE AN	ROTATIONS		Aity, Docket No. 44921-5039W0 Doc. No. 1793397.1
Seq. [D No.	(Iden(III)er I	Cenbenk Ace. Reil Seg. ID No.	Model Gode	Hondlogovs Cene Name	Homologous Gluster Name
NOS	<u>loenmer</u>	Ref-Seq-IP-NO.	Modelledde	dual-specificity tyrosine-(Y) phosphorylation regulated kinase 1A, dual-specificity tyrosine-(Y)- phosphorylation regulated	ESTs, Moderately similar to DYRK RAT DUAL-SPECIFICITY TYROSINE-(Y)-PHOSPHORYLATION REGULATED KINASE [R.norvegicus], ESTs, Weakly similar to DYRK MOUSE DUAL-SPECIFICITY TYROSINE-(Y)-PHOSPHORYLATION REGULATED KINASE [M.musculus], Homo sapiens chromosome 19, CIT-HSP BAC 470n8, Mus musculus, clone MGC:6699 IMAGE:3584001, mRNA, complete cds, dual-specificity tyrosine-(Y)-phosphorylation regulated kinase 1A, dual-specificity tyrosine-(Y)-phosphorylation regulated kinase 1a,
1540	18135	NM_012791	w	kinase 1a	kinase 2 GAMT HUMAN
				guanidinoacetate N- methyltransferase, quanidinoacetate	GUANIDINOACETATE N- METHYLTRANSFERASE [H.sapiens], Homo sapiens, clone MGC:14390 IMAGE:4300887, mRNA, complete cds, guanidinoacetate N- methyltransferase, guanidinoacetate
1541	16947	NM_012793	p,bb	methyltransferase	methyltransferase
1542	960	NM 012796		glutathione S-transferase theta 2, glutathione S-	ESTs, Highly similar to GTT2 RAT GLUTATHIONE S-TRANSFERASE YRS-YRS [R.norvegicus], Homo sapiens mRNA; cDNA DKFZp762N226 (from clone DKFZp762N226), expressed sequence Al266894, glutathione S-transferase theta 2, glutathione S-transferase, theta 2, but the time the set of the sequence Al266894.
			u	mal, T-cell differentiation protein, myelin and lymphocyte protein; T-cell	hypothetical protein BENE protein, Mus musculus, Similar to BENE protein, clone MGC:19097 IMAGE:4205488, mRNA, complete cds, RIKEN cDNA 2700018N07 gene, expressed sequence Al461653, mal, T- cell differentiation protein, mal, T-cell differentiation protein 2, myelin and lymphocyte protein; T-cell
1543	260	NM_012798	f,u	differentiation protein	differentiation protein B-factor, properdin, DKFZP586H2123
1544	556	NM_012803	d	protein C, protein C (inactivator of coagulation factors Va and VIIIa)	protein, ESTs, Weakly similar to PRTC RAT VITAMIN-K DEPENDENT PROTEIN C PRECURSOR [R.norvegicus], protein C, protein C (inactivator of coagulation factors Va and VIIIa)
1545	21729	NM 012804	q	ATP-binding cassette, sub- family D (ALD), member 3	ATP-binding cassette, sub-family D (ALD), member 1, ATP-binding cassette, sub-family D (ALD), member 2, ATP-binding cassette, sub-family D (ALD), member 3, ESTs, Weakly similar to ABD3 RAT ATP-BINDING CASSETTE, SUB-FAMILY D, MEMBER 3 [R.norvegicus]
				(ALD), Member 3	alpha-methylacyl-CoA racemase,
1546 1547	15032 24895	NM_012816 NM_012817	General General	insulin-like growth factor binding protein 5	hypothetical protein FLJ11808 insulin-like growth factor binding protein 5

TABLES	S IHWIMMAN	HOMOROGÚE AV	A SKIOITATIOKI		Atty: Docket No. 44921-5039WO Doc. No. 1793397.1
Seg. ID No.	lienWiler	Genbenk Ace./ Rai. Seq. ID No.	Model Gode	Homologous Cene Name	Homologovis Gluster Name
1548	18109	NM 012823	u,General		EST, Weakly similar to ANXA_HUMAN ANNEXIN XI [H.sapiens], annexin A10, annexin A3
				ATP-binding cassette, sub-	ATP-binding cassette, sub-family C (CFTR/MRP), member 10, ATP-binding cassette, sub-family C (CFTR/MRP), member 1a, ATP-binding cassette, sub-family C (CFTR/MRP), member 2, ATP-binding cassette, sub-family C (CFTR/MRP), member 4, ATP-binding cassette, sub-family C (CFTR/MRP), member 5, ESTS, Weakly similar to A40303 cystic fibrosis transmembrane conductance
1549	373	NM_012833	h,l,q,General	family C (CFTR/MRP), member 2	regulator - mouse [M.musculus], expressed sequence Al132311
1550	2855	NM_012838	e	cystatin B, cystatin B (stefin B)	cystatin B, cystatin B (stefin B), expressed sequence AA960480
				cytochrome c, cytochrome	ESTs, Highly similar to 630485A cytochrome c [H.sapiens], ESTs, Weakly similar to CYTOCHROME C, SOMATIC [M.musculus], Homo sapiens pseudogene for cytochrome clike protein, clone pHGC4E1, Human DNA sequence from clone RP11-169017 on chromosome 13 Contains ESTs, GSSs, STSs and four CpG islands. Contains a novel protein similar to cytochrome c, part of a novel gene similar to TPTE encoding a transmembrane phosphatase with tensin homology and the ADPRTL1 gene encoding ADP-ribosyltransferase (NAD+; poly (ADP-ribose) polymerase)-like protein 1 (vault protein, KIAA0177), cytochrome c, cytochrome
1551	11136	NM_012839	s	c, somatic	c, somatic EST, Moderately similar to
1552	20885	NM 012842	а	epidermal growth factor, epidermal growth factor (beta-urogastrone)	EPIDERMAL GROWTH FACTOR PRECURSOR [M.musculus], ESTs, Moderately similar to EPIDERMAL GROWTH FACTOR PRECURSOR [M.musculus], ESTs, Weakly similar to EGRT epidermal growth factor precursor - rat [R.norvegicus], Homo sapiens mRNA; cDNA DKFZp434O0213 (from clone DKFZp434O0213); partial cds, epidermal growth factor, epidermal growth factor (beta-urogastrone), nidogen 2
1552	20884	NM_012842	a,bb	epidermal growth factor, epidermal growth factor (beta-urogastrone)	EST, Moderately similar to EPIDERMAL GROWTH FACTOR PRECURSOR [M.musculus], ESTs, Moderately similar to EPIDERMAL GROWTH FACTOR PRECURSOR [M.musculus], ESTs, Weakly similar to EGRT epidermal growth factor precursor - rat [R.norvegicus], Homo sapiens mRNA; cDNA DKFZp434O0213 (from clone DKFZp434O0213); partial cds, epidermal growth factor, epidermal growth factor (beta-urogastrone), nidogen 2

TABLE	SE HÜMAN	HOWO FOCA NEW	ROTESTON		Aity. Docket No. 44921-5089W0 Doc. No. 1793897.1
Seq. ID No.	ldentifier	Condent Ace. Ref. See. ID No.	Model Code	Homologous Cene Name	Homologous Gluster Name
1553	18770	NM_012857	е	lysosomal membrane glycoprotein 1, lysosomal- associated membrane protein 1	CD68 antigen, ESTs, Weakly similar to LMP1 RAT LYSOSOME-ASSOCIATED MEMBRANE GLYCOPROTEIN 1 PRECURSOR [R.norvegicus], lysosomal membrane glycoprotein 1, lysosomal-associated membrane protein 1, similar to \$68401 (cattle) glucose induced gene
1554	20674	NM_012861	ì	0-6-methylguanine-DNA methyltransferase, O-6- methylguanine-DNA methyltransferase	ESTs, Highly similar to 1207289A reverse transcriptase related protein [H.sapiens], ESTs, Highly similar to S21348 probable pol polyprotein-related protein 4 - rat [R.norvegicus], ESTs, Moderately similar to GNMSLL retrovirus-related reverse transcriptase homolog - mouse retrotransposon [M.musculus], ESTs, Weakly similar to 1207289A reverse transcriptase related protein [H.sapiens], ESTs, Weakly similar to B34087 hypothetical protein [H.sapiens], Homo sapiens cDNA FLJ12202 fis, clone MAMMA1000908, Mus musculus, Similar to L1 repeat, Tf subfamily, member 30, clone MGC:7372 IMAGE:3487559, mRNA, complete cds, RIKEN cDNA 1700082M22 gene, T lymphoma oncogene, expressed sequence Al267024, hypothetical protein FLJ21032, pheromone receptor V3R4
1555	13151	NM_012862	a,r,General	matrix Gla protein, matrix gamma-carboxyglutamate (gla) protein	MATRIX GLA-PROTEIN PRECURSOR [H.sapiens], matrix Gla protein, matrix gamma- carboxyglutamate (gla) protein
1556	24617	NM_012870	General	tumor necrosis factor receptor superfamily, member 11b (osteoprotegerin)	Nerve growth factor receptor, fast, RIKEN cDNA 2610311B09 gene, nerve growth factor receptor, nerve growth factor receptor, nerve growth factor receptor (TNFR superfamily, member 16), tumor necrosis factor receptor superfamily, member 11b (osteoprotegerin), tumor necrosis factor receptor superfamily, member 21 EST, Moderately similar to 60S RIBOSOMAL PROTEIN L39 [R.norvegicus], ESTs, Highly similar to 60S RIBOSOMAL PROTEIN L39 [R.norvegicus], ESTs, Highly similar to G02654 ribosomal protein L39 [H.sapiens], ESTs, Moderately similar to G02654 ribosomal protein L39 [H.sapiens], RIKEN cDNA 2810465O16 gene, RIKEN cDNA
1557	20945	NM_012875	a,v		4930517K11 gene, ribosomal protein L39, ribosomal protein L39-like 1

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TABLE	BA HUMAN	HONOLOGUEA	<u>enonatōni</u>		Any, Docket No. 44921-5039WO Doc. No. 1799597.1
Seq. (D No.	ldenilijer	Confinit Aced Ref. Seq. ID No.	Modal Goda	Homologous Gano Namo	Homologous Gluster Nams
1558	15872	NM_012879	o,r	solute carrier family 2 (facilitated glucose transporter), member 2 superoxide dismutase 3,	ESTs, Highly similar to A31318 glucose transporter-like protein [H.sapiens], ESTs, Weakly similar to GLUCOSE TRANSPORTER TYPE 2, LIVER [R.norvegicus], ESTs, Weakly similar to S05319 glucose transport protein, hepatic - mouse [M.musculus], solute carrier family 2 (facilitated glucose transporter), member 10, solute carrier family 2 (facilitated glucose transporter), member 2, solute carrier family 2, (facilitated glucose transporter) member 8 RIKEN cDNA 1700105P06 gene,
1559	495	NM_012880	z	extracellular superoxide dismutase 3,	superoxide dismutase 3, extracellular RIKEN cDNA 1700105P06 gene.
1559	494	NM_012880	С	extracellular	superoxide dismutase 3, extracellular
1560	23651	NM 012881	d,u,General	secreted phosphoprotein 1, secreted phosphoprotein 1 (osteopontin, bone sialoprotein I, early T-lymphocyte activation 1)	
1562	19477	NM_012891	q		
1563	18564	NM_012899	v,General	aminolevulinate, delta-, dehydratase	aminolevulinate, delta-, dehydratase
1564 1564	7197 7196	NM_012904 NM_012904	f,r,cc,General	annexin A1	EST, Weakly similar to A Chain A, Nmr Solution Structure Of Domain 1 Of Human Annexin I (SUB 41-113 [H.sapiens], annexin A1
1565	20202	NM_012909	b,r	aquaporin 2, aquaporin 2 (collecting duct)	aquaporin 2, aquaporin 2 (collecting duct), aquaporin 6, aquaporin 6, kidney specific
1566	16581	NM_012911	c,j		Mus musculus retinal cone arrestin 3 (Arr3) mRNA, complete cds, RIKEN cDNA 1200006117 gene, arrestin, beta 2, expressed sequence Al326910, retinal S-antigen Mus musculus retinal cone arrestin 3
1566	16582	NM_012911	c		(Arr3) mRNA, complete cds, RIKEN cDNA 1200006117 gene, arrestin, beta 2, expressed sequence Al326910, retinal S-antigen
1567	24431	NM_012912	General	activating transcription factor 3	ESTs, Highly similar to 1604249C transcription factor ATF3 [H.sapiens], ESTs, Weakly similar to ATF3 RAT CYCLIC-AMP-DEPENDENT TRANSCRIPTION FACTOR ATF-3 [R.norvegicus], Jun dimerization protein 2, activating transcription factor 3, basic leucine zipper transcription factor, ATF-like
1568	18118	NM_012913	p	ATPase, Na+/K+ transporting, beta 3 polypeptide	ATPase, Na+/K+ transporting, beta 3 polypeptide, ESTs, Highly similar to G02485 Na+/K+-exchanging ATPase [H.sapiens], expressed sequence Al664000
1569	6108	NM_012915	n	polypopude	ATPase inhibitor, ATPase inhibitor precursor, ESTs, Moderately similar to ATPASE INHIBITOR, MITOCHONDRIAL PRECURSOR [R.norvegicus], Mus musculus 10 days embryo cDNA, RIKEN full-length enriched library, clone:2610204M17, full insert sequence

TABLES	E HUMAN	HOMOROGAE VA	INOTATIONS		Aitty. Docket No. 4/1921-5039WO Doc. No. 1793897.1
(2007. [D) (108	ldenililer	CenBank Acc./ Ref. Seq. ID No.	Model Gode	Homologous Cono Name	Homologous Gluster Name
1570	20757	NM_012923	c,i,aa	cyclin G, cyclin G1	cyclin G, cyclin G1, cyclin G2, cyclin I
1570	20755	NM_012923	i	cyclin G, cyclin G1	cyclin G, cyclin G1, cyclin G2, cyclin I
1571	2830	NM_012925	f	CD59 antigen p18-20 (antigen identified by monoclonal antibodies 16.3A5, EJ16, EJ30, EL32 and G344), CD59a antigen	CD59 antigen p18-20 (antigen identified by monoclonal antibodies 16.3A5, EJ16, EJ30, EL32 and G344), CD59a antigen, ESTs, Weakly similar to CD59 RAT CD59 GLYCOPROTEIN PRECURSOR [R.norvegicus]
1571	2831	NM_012925	f	CD59 antigen p18-20 (antigen identified by monoclonal antibodies 16.3A5, EJ16, EJ30, EL32 and G344), CD59a antigen	CD59 antigen p18-20 (antigen identified by monoclonal antibodies 16.3A5, EJ16, EJ30, EL32 and G344), CD59a antigen, ESTs, Weakly similar to CD59 RAT CD59 GLYCOPROTEIN PRECURSOR [R.norvegicus]
1572	1977	NM_012930	q	carnitine palmitoyltransferase 2, carnitine palmitoyltransferase II	EST, Moderately similar to CPT2 RAT CARNITINE O-PALMITOYLTRANSFERASE II, MITOCHONDRIAL PRECURSOR [R.norvegicus], carnitine palmitoyltransferase 2, carnitine palmitoyltransferase II, expressed sequence Al323697
1573	18694	NM_012931	j.l,m,z		CD2-associated protein, HEF like Protein, Homo sapiens cDNA FLJ14854 fis, clone PLACE1000972, breast cancer anti-estrogen resistance 1, enhancer of filamentation 1 (cas-like docking; Crk-associated substrate related), neural precursor cell expressed, developmentally downregulated gene 9, v-crk-associated tyrosine kinase substrate
1574	13723	NM_012935	n	crystallin, alpha B	
1575 1575	9109 19398	NM_012939 NM_012939	j,y,z aa	cathepsin H	ESTs, Highly similar to KHHUH cathepsin H [H.sapiens], ESTs, Weakly similar to CATHEPSIN H PRECURSOR [R.norvegicus], Homo sapiens cDNA: FLJ22499 fis, clone HRC11250, highly similar to HSCATHH Human mRNA for cathepsin H (EC 3.4.22.16), cathepsin H, cathepsin W, cathepsin W (lymphopain)
				diphtheria toxin receptor	
1576	223	NM_012945	b,cc		diphtheria toxin receptor (heparin- binding epidermal growth factor-like growth factor), expressed sequence AW047313, heparin binding epidermal growth factor-like growth factor ESTs, Weakly similar to
1577	15058	NM_012950	cc	coagulation factor II (thrombin) receptor	GPRY_MOUSE PROBABLE G PROTEIN-COUPLED RECEPTOR GPR34 [M.musculus], G protein- coupled receptor 41, G protein- coupled receptor 43, Rattus norvegicus protease activated receptor 3 mRNA, complete cds, coagulation factor II (thrombin) receptor

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		HOMOFOGAEV	SKOLĽŠLOKI		Affy, Docket No. 44921-5039W0 Doc. No. 1798397.1
Seq. ID No.	ldentifier	GenBenk Aced Red Seq. ID No. '	Model Code	Kama Kamalogove Gene	Homologous Cluster Name
				high mobility group box 1, high-mobility group (nonhistone chromosomal)	EST, Moderately similar to A Chain A, Crystal Structure Of Hmg1 Domain A Bound To A Cisplatin- Modified Dna Duplex [R.norvegicus], EST, Weakly similar to A Chain A, Crystal Structure Of Hmg1 Domain A Bound To A Cisplatin- Modified Dna Duplex [R.norvegicus], ESTs, Highly similar to S02826 nonhistone chromosomal protein HMG-1 [H.sapiens], ESTs, Moderately similar to HIGH MOBILITY GROUP PROTEIN HMG1 [M.musculus], RIKEN cDNA 4932431P20 gene, high mobility group box 1, high mobility group box 3, highmobility group (nonhistone chromosomal) protein 1, high-mobility group (nonhistone chromosomal) protein 1-like 10, high-mobility group (nonhistone chromosomal) protein 1-
1579	19111	NM_012963	g	protein 1	like 3 Mus musculus 12 days embryo male
1580	19374	NM 012964	x	hyaluronan mediated motility receptor (RHAMM), hyaluronan-mediated motility receptor (RHAMM)	wolffian duct includes surrounding region cDNA, RIKEN full-length enriched library, clone:6720466F14, full insert sequence, RIKEN cDNA 0610027D24 gene, TRAF4 associated factor 1, hyaluronan mediated motility receptor (RHAMM), hyaluronan-mediated motility receptor (RHAMM)
1581	2554	NM_012967	t	intercellular adhesion molecule, intercellular adhesion molecule 1 (CD54), human rhinovirus receptor	Homo sapiens mRNA; cDNA DKFZp434E0516 (from clone DKFZp434E0516), intercellular adhesion molecule, intercellular adhesion molecule 1 (CD54), human rhinovirus receptor, intercellular adhesion molecule 3, intercellular adhesion molecule 5, telencephalin
1581	2555	NM_012967	t,cc,General	intercellular adhesion molecule, intercellular adhesion molecule 1 (CD54), human rhinovirus receptor	Homo sapiens mRNA; cDNA DKFZp434E0516 (from clone DKFZp434E0516), intercellular adhesion molecule, intercellular adhesion molecule 1 (CD54), human rhinovirus receptor, intercellular adhesion molecule 3, intercellular adhesion molecule 5, telencephalin
1582	24528	NM 012973	С	potassium voltage-gated channel, Isk-related family, member 1, potassium voltage-gated channel, Isk- related subfamily, member 1	potassium voltage-gated channel, Isk- related family, member 1, potassium voltage-gated channel, Isk-related subfamily, member 1
1583	956	NM_012976	c		ESTs, Highly similar to LEG9 RAT GALECTIN-9 [R.norvegicus], ESTs, Highly similar to LEG9_HUMAN GALECTIN-9 [H.sapiens], ESTs, Weakly similar to LEG9 RAT GALECTIN-9 [R.norvegicus], expressed sequence Al265545, lectin, galactose binding, soluble 12, lectin, galactose binding, soluble 9, lectin, galactoside-binding, soluble, 9 (galectin 9)
1584	16417	NM_012991	g	nucleoporin 50kD, nucleoprotein 50	nucleoporin 50kD, nucleoprotein 50

TABLE	MAMUH S	HOWOLOGAE VA	ROTATIONS		Ally, Docket No. 44921-5039WC Doc. No. 1793397.
Seg. [D] No.	ldentifer	Cenbenk Acc./ Ref. Seq. ID No.	Model Code 4	Homologous Cene Name	Homologous Gluster Namo
1585	17393	NM_012992	d	nucleophosmin (nucleolar phosphoprotein B23, numatrin), nucleophosmin 1	ESTs, Moderately similar to NPM_HUMAN NUCLEOPHOSMIN [H.sapiens], ESTs, Weakly similar to NPM_HUMAN NUCLEOPHOSMIN [H.sapiens], nucleophosmin (nucleolar phosphoprotein B23, numatrin), nucleophosmin 1, nucleophosmin/nucleoplasmin 3, nucleoplasmin 3
1586	23544	NM 013013	s	prosaposin, prosaposin (variant Gaucher disease and variant metachromatic leukodystrophy)	ESTs, Weakly similar to 1504251A sphingolipid activator [H.sapiens], RIKEN cDNA 2310020A21 gene, prosaposin, prosaposin (variant Gaucher disease and variant metachromatic leukodystrophy)
1587	1588	NM 013026	k	syndecan 1	syndecan 1
1588	17894	NM 013027	m	selenoprotein W, 1,	ESTs, Weakly similar to SELW MOUSE SELENOPROTEIN W [M.musculus], selenoprotein W, 1, selenoprotein W, muscle 1
1590	18300	NIM 013020	a v Conoral		ESTs, Weakly similar to NPT2 RAT RENAL SODIUM-DEPENDENT PHOSPHATE TRANSPORT PROTEIN 2 [R.norvegicus], ESTs, Weakly similar to NPT2_HUMAN RENAL SODIUM-DEPENDENT PHOSPHATE TRANSPORT PROTEIN 2 [H.sapiens], Homo sapiens, Similar to solute carrier family 34 (sodium phosphate), member 1, clone MGC:18179 IMAGE:4155326, mRNA, complete cds, Solute carrier family 17 (sodium/hydrogen exchanger), member 2, expressed sequence Al649385, solute carrier family 34 (sodium phosphate), member 3, solute carrier family 34 (sodium phosphate), member 3
1589	18300	NM_013030	s,v,General		phosphate), member 2 ESTs, Weakly similar to NPT2 RAT RENAL SODIUM-DEPENDENT PHOSPHATE TRANSPORT PROTEIN 2 [R.norvegicus], ESTs, Weakly similar to NPT2_HUMAN RENAL SODIUM- DEPENDENT PHOSPHATE TRANSPORT PROTEIN 2 [H.sapiens], Homo sapiens, Similar to solute carrier family 34 (sodium phosphate), member 1, clone MGC:18179 IMAGE:4155326, mRNA, complete cds, Rattus norvegicus mRNA for NaPi-2 alpha, complete cds, expressed sequence
				solute carrier family 34 (sodium phosphate).	Al649385, solute carrier family 34 (sodium phosphate), member 1, solute carrier family 34 (sodium phosphate),

TABLES	HUMAN	HOMOROGUEAN	RADITATIONS		Aity, Dooket No. 44921-5039WO Doc. No. 1793397.1
Seq.(D)		ConBank Aced Ref. Seq. ID No.	Madel Code	Homologous Cene): Name	Homologous Cluster Name
				solute carrier family 34 (sodium phosphate),	ESTs, Weakly similar to NPT2 RAT RENAL SODIUM-DEPENDENT PHOSPHATE TRANSPORT PROTEIN 2 [R.norvegicus], ESTs, Weakly similar to NPT2_HUMAN RENAL SODIUM-DEPENDENT PHOSPHATE TRANSPORT PROTEIN 2 [H.sapiens], Homo sapiens, Similar to solute carrier family 34 (sodium phosphate), member 1, clone MGC:18179 IMAGE:4155326, mRNA, complete cds, Rattus norvegicus mRNA for NaPi-2 alpha, complete cds, Solute carrier family 17 (sodium/hydrogen exchanger), member 2, expressed sequence Al649385, solute carrier family 34 (sodium phosphate), member 1, solute carrier family 34 (sodium phosphate),
1589	18078	NM_013030	s	member 1	member 2 ESTs, Weakly similar to NPT2 RAT
1589	18077	NM_013030	e,s,z	solute carrier family 34 (sodium phosphate), member 1	RENAL SODIUM-DEPENDENT PHOSPHATE TRANSPORT PROTEIN 2 [R.norvegicus], ESTs, Weakly similar to NPT2_HUMAN RENAL SODIUM- DEPENDENT PHOSPHATE TRANSPORT PROTEIN 2 [H.sapiens], Homo sapiens, Similar to solute carrier family 34 (sodium phosphate), member 1, clone MGC:18179 IMAGE:4155326, mRNA, complete cds, Rattus norvegicus mRNA for NaPi-2 alpha, complete cds, expressed sequence Al649385, solute carrier family 34 (sodium phosphate), member 1, solute carrier family 34 (sodium phosphate), member 2
1591	730	NM_013040	w	ATP-binding cassette, sub- family C (CFTR/MRP), member 9	ATP-binding cassette, sub-family C (CFTR/MRP), member 8, ATP-binding cassette, sub-family C (CFTR/MRP), member 9, ESTs, Highly similar to ACC8_HUMAN SULFONYLUREA RECEPTOR 1 [H.sapiens], ESTs, Weakly similar to T42728 sulfonylurea receptor 2, isoform B - mouse [M.musculus], Mus musculus adult male pituitary gland cDNA, RIKEN full-length enriched library, clone:5330439B14, full insert sequence
1592	17401	NM_013043	i,o,General	transforming growth factor beta 1 induced transcript 4, transforming growth factor beta-stimulated protein TSC-22	ESTs, Moderately similar to DIP_HUMAN DIP PROTEIN [H.sapiens], delta sleep inducing peptide, immunoreactor, glucocorticoid-induced leucine zipper, transforming growth factor beta 1 induced transcript 4, transforming growth factor beta-stimulated protein TSC-22
1593	16684	NM_013052	General	tyrosine 3- monooxygenase/tryptopha n 5-monooxygenase activation protein, eta polypeptide	3-monooxgenase/tryptophan 5- monooxgenase activation protein, gamma polypeptide, tyrosine 3- monooxygenase/tryptophan 5- monooxygenase activation protein, eta polypeptide

TRABILE S	KAMUKI S	HOMOLOGNE V	INOTATIONS		A11y., Docket No. 44921-5039W0 Doc. No. 1793397.1
Seq. ID No.	ldeniiier	Consent Ace./ Ref. Seq. ID No.	Model Code	Homologous Cono Namo	Homologous Gluster Name
					RIKEN cDNA 2700028P07 gene, tyrosine 3-monooxygenase/tryptophan 5-monooxygenase activation protein,
1594	14421	NM_013053	u		theta polypeptide inhibitor of DNA binding 3, inhibitor of
1595	15254	NM_013058	k		DNA binding 3, dominant negative helix-loop-helix protein alkaline phosphatase,
1596	14997	NM_013059	s,z		liver/bone/kidney alkaline phosphatase,
1596 1597	14996 25676	NM_013059 NM_013069	General aa		liver/bone/kidney
					CD74 antigen (invariant polypeptide of major histocompatibility complex, class II antigen-associated), la-associated invariant chain, KIAA0275 gene product, sparc/osteonectin, cwcv and kazal-like domains proteoglycan (testican), sparc/osteonectin, cwcv and kazal-like domains proteoglycan 1, sparc/osteonectin, cwcv and kazal-like
1597	16924	NM_013069	0		domains proteoglycan 2 Dystrophin, dystrophin (muscular dystrophy, Duchenne and Becker types), includes DXS142, DXS164, DXS206, DXS230, DXS239, DXS268, DXS269, DXS270, DXS272, dystrophin, muscular dystrophy, dystrophin-related protein 2 A-form
1598	24748	NM_013070	h,q	utrophin, utrophin (homologous to dystrophin)	splice variant, utrophin, utrophin (homologous to dystrophin)
1599	1529	NM_013082	d,General		syndecan 2, syndecan 2 (heparan sulfate proteoglycan 1, cell surface- associated, fibroglycan)
1600	1521	NM_013091	j, l ,z,General		tumor necrosis factor receptor superfamily, member 12, tumor necrosis factor receptor superfamily, member 12 (translocating chainassociation membrane protein), tumor necrosis factor receptor superfamily, member 1A, tumor necrosis factor receptor superfamily, member 1a, tumor necrosis factor receptor superfamily, member 1a-like 1, tumor necrosis factor receptor superfamily, member 1a-like 1, tumor necrosis factor receptor superfamily, member 1a-like 2
			,,,,,,	hemoglobin alpha, adult chain 2, hemoglobin, alpha	
1601	1685	NM_013096	c,aa	1	
1601	26150	NM_013096	c,i	hemoglobin alpha, adult	
1601	1688	NM_013096	р	chain 2, hemoglobin, alpha	
1601	1689	NM 013096	c,p	hemoglobin alpha, adult chain 2, hemoglobin, alpha	EST, Moderately similar to HART1 hemoglobin alpha-1 chain - rat [R.norvegicus], ESTs, Moderately similar to HART1 hemoglobin alpha-1 chain - rat [R.norvegicus], RIKEN cDNA 2510042H12 gene, hemoglobin alpha, adult chain 1, hemoglobin, alpha 1, hemoglobin, alpha 2

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TABLE	KAMUH,	HONOTOGAE VY	INOTATIONS		Ally, Docket No. 44921-5039WC Doc. No. 1793397.1
Seq. ID : No.	Identifier	CenBenk Ace./ Ref. Seq. ID No.:	Model Code:	Homologous Cene Name	Homologovis Clyster Name
1601 1602	1684 20886	NM_013096 NM_013097	c,s,aa u,x,bb	hemoglobin alpha, adult chain 2, hemoglobin, alpha 1	EST, Moderately similar to HART1 hemoglobin alpha-1 chain - rat [R.norvegicus], ESTs, Moderately similar to HART1 hemoglobin alpha-1 chain - rat [R.norvegicus], RIKEN cDNA 2510042H12 gene, hemoglobin alpha, adult chain 1, hemoglobin, alpha 1, hemoglobin, alpha 2
1002	20000	1410 013097	u,x,bb		ESTs, Weakly similar to DRN1 RAT
1602	20887	NM_013097	u,×,bb		DEOXYRIBONUCLEASE I PRECURSOR [R.norvegicus], Mus musculus DNasel precursor mRNA, complete cds, RIKEN cDNA 4733401H14 gene, deoxyribonuclease I, expressed sequence Al788650 ESTs, Weakly similar to G6PT RAT
1603	1321	NM_013098	С		GLUCOSE-6-PHOSPHATASE [R.norvegicus], Homo sapiens, clone IMAGE:3050476, mRNA, partial cds, RIKEN cDNA 0710001K01 gene, expressed sequence AW545836, glucose-6-phosphatase, catalytic, glucose-6-phosphatase, catalytic (glycogen storage disease type I, von Gierke disease), glucose-6- phosphatase, catalytic, related sequence, islet-specific glucose-6- phosphatase catalytic subunit-related protein
1604	15296	NM_013102	l,m	FK506 binding protein 1a (12 kDa), FK506-binding protein 1A (12kD)	ESTs, Moderately similar to 1613455A FK506 binding protein FKBP [H.sapiens], FK506 binding protein 1a (12 kDa), FK506 binding protein 1b (12.6 kDa), FK506 binding protein 2 (13 kDa), FK506 binding protein 4 (59 kDa), FK506 binding protein 5 (51 kDa), FK506-binding protein 1A (12kD)
1606	22700	NIM 013113	0.07.00	_	ATPase, Na+/K+ transporting, beta 1
1606	23709	NM_013113 NM_013113	p		polypeptide ATPase, Na+/K+ transporting, beta 1 polypeptide
1606	23710	NM_013113	s		ATPase, Na+/K+ transporting, beta 1 polypeptide
1607	1976	NM_013118	u		guanylate cyclase activator 1B (retina), guanylate cyclase activator 2 (guanylin 2, intestinal, heatstable), guanylate cyclase activator 2A (guanylin)
1609	870	NM_013130	h		MAD (mothers against decapentaplegic, Drosophila) homolog 1, MAD (mothers against decapentaplegic, Drosophila) homolog 5, MAD (mothers against decapentaplegic, Drosophila) homolog 9
1610	16650	NM_013132	u,General	annexin A5	annexin A5
1611	650	NM_013134	h		3-hydroxy-3-methylglutaryl-Coenzyme A reductase, ESTs, Moderately similar to hydroxymethylglutaryl-CoA reductase [M.musculus], SREBP CLEAVAGE-ACTIVATING PROTEIN

TABLE	e Human	HOMOLOGUE A	SKOTFATOKI	Ant.	i Airy, Docker (10, 44921-5039)(0
Sea. ID		Censenk Acc./		Homologous Gino H	<u>Doc. No. 1793397.1</u>
No.	Identifier	Ref. Seq. ID No.		Name :	Homologous Gluster Name
					3-hydroxy-3-methylglutaryl-Coenzyme
					A reductase, ESTs, Moderately similar to hydroxymethylglutaryl-CoA
					reductase [M.musculus], SREBP
1611	651	NM_013134	h,j,l		CLEAVAGE-ACTIVATING PROTEIN
					ESTs, Moderately similar to INOSITOL
					1,4,5-TRISPHOSPHATE-BINDING
					PROTEIN TYPE 1 RECEPTOR [M.musculus], Mus musculus, Similar
					to Purkinje cell protein 1, clone
					MGC:11943 IMAGE:3600031, mRNA,
					complete cds, RIKEN cDNA
				inositol 1,4,5-triphosphate	9330127I20 gene, inositol 1,4,5- triphosphate receptor 1, inositol 1,4,5-
				receptor 3, inositol 1,4,5-	triphosphate receptor 2, inositol 1,4,5-
					triphosphate receptor 5, inositol 1,4,5-
1612	1712	NM_013138	General	3	triphosphate receptor, type 3
					insulin-like growth factor binding
1613	16982	NM_013144	o,v,General		protein 1, protease, serine, 11 (IGF binding)
1013	10302	1411_013144	U,V,Oerierat	CCAAT/enhancer binding	CCAAT/enhancer binding protein
1614	21683	NM_013154	t,cc,General	protein (C/EBP), delta	(C/EBP), delta
				CCAAT/enhancer binding	CCAAT/enhancer binding protein
1614	21682	NM_013154	СС	protein (C/EBP), delta	(C/EBP), delta
					ESTs, Weakly similar to CATHEPSIN L PRECURSOR [M.musculus], RIKEN
					cDNA 2310051M13 gene, RIKEN
					cDNA 4930486L24 gene, cathepsin 7,
1015		040450			cathepsin F, cathepsin L, cathepsin
1615 1615	3431 25567	NM_013156 NM_013156	b,g,n v,General	cathepsin L	L2, cathepsin O
1013_	20007	14M_010100	v,ocneral		ESTs, Weakly similar to CATHEPSIN
					L PRECURSOR [M.musculus], RIKEN
					cDNA 2310051M13 gene, RIKEN
					cDNA 4930486L24 gene, cathepsin 7, cathepsin F, cathepsin L, cathepsin
1615	3430	NM_013156	General	cathepsin L	L2, cathepsin O
					RIKEN cDNA 4833415K22 gene,
					expressed sequence AA675336,
1616	1309	NM 013159	l _w		insulin degrading enzyme, insulin- degrading enzyme
1010	1309	1414 013139	W		RIKEN cDNA 4833415K22 gene,
					expressed sequence AA675336,
					insulin degrading enzyme, insulin-
1616	1310	NM_013159	w		degrading enzyme
					growth factor, beta 2, transforming
1617	21723	NM_013174	l _w		growth factor, beta 3
					EST, Moderately similar to CAMP-
					DEPENDENT PROTEIN KINASE
					TYPE I-ALPHA REGULATORY CHAIN
					[R.norvegicus], protein kinase, cAMP dependent regulatory, type 1, alpha,
					protein kinase, cAMP dependent
					regulatory, type I beta, protein kinase,
4040		NINA 040404	<u> </u>		cAMP-dependent, regulatory, type I,
1618	1314	NM_013181	lm	meprin 1 beta, meprin A,	alpha (tissue specific extinguisher 1) expressed sequence C87576, meprin
1619	17357	NM 013183	p,bb,General	beta	1 beta, meprin A, beta
	T				Mus musculus adult male stomach
					cDNA, RIKEN full-length enriched
					library, clone:2210403E17, full insert
					sequence, expressed sequence AA407869, phosphofructokinase, liver,
					phosphofructokinase, liver, B-type,
1620	1300	NM_013190	ly	1	phosphofructokinase, muscle

TABLE 8	e HUMAN	HOMO FOGUE VA	ZNOTKATONI		Airy, Docket No. 44921-5039Wo Dock No. 1793397.1
Seq. ID: No:	(dentifier	GenBenk Aced Ref. Seq. ID No.	Model Gode	Homológous Cens Nems	Homologous Cluster Name
1621	16448	NM_013197	С		EST, Highly similar to HEM0 RAT 5-AMINOLEVULINIC ACID SYNTHASE, ERYTHROID-SPECIFIC, MITOCHONDRIAL PRECURSOR [R.norvegicus], ESTs, Highly similar to SYHUAE 5-aminolevulinate synthase [H.sapiens], aminolevulinate, delta-, synthase 2 (sideroblastic/hypochromic anemia), aminolevulinic acid synthase 1, aminolevulinic acid synthase 2, erythroid, glycine C-acetyltransferase (2-amino-3-ketobutyrate coenzyme A ligase), glycine C-acetyltransferase (2-amino-3-ketobutyrate-coenzyme A ligase)
1622	20856	NM_013200	b	carnitine palmitoyltransferase 1, muscle, carnitine palmitoyltransferase I, muscle	ESTs, Moderately similar to CPTM RAT CARNITINE O- PALMITOYLTRANSFERASE I, MITOCHONDRIAL MUSCLE ISOFORM [R.norvegicus], ESTs, Weakly similar to CPT1 MOUSE CARNITINE O- PALMITOYLTRANSFERASE I, MITOCHONDRIAL LIVER ISOFORM [M.musculus], carnitine palmitoyltransferase 1, liver, carnitine palmitoyltransferase 1, muscle, carnitine palmitoyltransferase I, muscle
1623	397	NM_013214	f		COTO LIGHT SINGLE ADZO LINAAN
					ESTs, Highly similar to AR72_HUMAN AFLATOXIN B1 ALDEHYDE REDUCTASE 1 (AFB1-AR 1) (ALDOKETOREDUCTASE 7) [H.sapiens], ESTs, Moderately similar to AFAR RAT AFLATOXIN B1 ALDEHYDE REDUCTASE [R.norvegicus], RIKEN cDNA 0610025K21 gene, Rattus norvegicus aiar mRNA for androgen-inducible aldehyde reductase, complete cds, aldo-keto reductase family 7, member A2 (aflatoxin aldehyde reductase), aldo-keto reductase family 7, member
1624	20864	NM_013215	g,n,y	- with believe of the control of the	A3 (aflatoxin aldehyde reductase) ESTs, Moderately similar to T30989
1625	20728	NM_013217	V		es is, Moderately similar to 130989 serine/threonine protein kinase NIK - mouse [M.musculus], Mus musculus, Similar to zinc finger protein 347, clone MGC:18913 IMAGE:4242025, mRNA, complete cds, RIKEN cDNA 1500031A17 gene, mitogen-activated protein kinase kinase kinase kinase kinase kinase kinase kinase kinase kinase kinase kinase kinase kinase kinase (green) (liping) (
		-	2	growth factor, erv1 (S.	ESTs, Highly similar to ALR RAT AUGMENTER OF LIVER REGENERATION [R.norvegicus],
1626	1396	NM_013222			growth factor, erv1 (S. cerevisiae)-like (augmenter of liver regeneration)
للبعين		010222		or made regendration)	(augmenter of fiver regeneration)

TABLES	e HUMAN	HOMOLOGUE AN	INOTATIONS		Atty. Docket No. 44921-5039W0 Doc. No. 1793397.1
Seq. [D No-	ldentifier	ConBent Acel Ref. Sog. D. No.		Homologous Cene Neme 1 4	Homologovis Cluster Name
1627	815	NM 013224		Before and a house of the property of the prop	ESTs, Highly similar to RS26_HUMAN 40S RIBOSOMAL PROTEIN S26 [H.sapiens], Homo sapiens, clone IMAGE:4100953, mRNA, Human DNA sequence from PAC 384D21 on chromosome X contains ribosomal protein S26 pseudogene, STS, polymerase (RNA) II (DNA directed) polypeptide D, ribosomal protein S26
1628	18305	NM_013226	v		
1629	21078	NM_016986	d	acetyl-Coenzyme A dehydrogenase, medium chain, acyl-Coenzyme A dehydrogenase, C-4 to C- 12 straight chain	ESTs, Highly similar to ACDM MOUSE ACYL-COA DEHYDROGENASE, MEDIUM-CHAIN SPECIFIC PRECURSOR [M.musculus], acetyl-Coenzyme A dehydrogenase, medium chain, acyl-Coenzyme A dehydrogenase, C-4 to C-12 straight chain, expressed sequence Al987948 ESTs, Weakly similar to PPAL RAT LYSOSOMAL ACID PHOSPHATASE
1630	24649	NM_016988	v	acid phosphatase 2, lysosomal	PRECURSOR [R.norvegicus], acid phosphatase 2, lysosomal, acid phosphatase 6, lysophosphatidic, acid phosphatase, prostate, acid phosphatase, testicular EST, Weakly similar to RL15 RAT 60S
1631	15239	NM_016989	q,w		RIBOSOMAL PROTEIN L15 [R.norvegicus], ESTs, Highly similar to RL15_HUMAN 60S RIBOSOMAL PROTEIN L15 [H.sapiens], ESTs, Moderately similar to RL15 RAT 60S RIBOSOMAL PROTEIN L15 [R.norvegicus], RIKEN cDNA 2510008H07 gene, ribosomal protein L15
1632	45	NM 016996	General	calcium-sensing receptor, calcium-sensing receptor (hypocalciuric hypercalcemia 1, severe neonatal hyperparathyroidism)	EST, Weakly similar to EXTRACELLULAR CALCIUM- SENSING RECEPTOR PRECURSOR [R.norvegicus], ESTs, Weakly similar to CASR_HUMAN EXTRACELLULAR CALCIUM-SENSING RECEPTOR PRECURSOR [H.sapiens], ESTs, Weakly similar to JC7160 metabotropic glutamate receptor subtype 3 precursor - mouse [M.musculus], G protein coupled receptor, family C, group 1, member A, G protein coupled receptor, family C, group 1, member C, calcium-sensing receptor (hypocalciuric hypercalcemia 1, severe neonatal hyperparathyroidism), vomeronasal 2, receptor, 11, vomeronasal 2, receptor, 12
1032	45	ININ_0.10990	General	cytochrome P450,	Mus musculus, Similar to cytochrome P450, 4a10, clone MGC:25972 IMAGE:4240359, mRNA, complete
1633	20714	NM_016999	t	subfamily IV B, polypeptide 1, cytochrome P450, subfamily IVB, polypeptide 1	cds, RIKEN cDNA A230105L22 gene, cytochrome P450, 4a10, cytochrome P450, 4a14, cytochrome P450, subfamily IVA, polypeptide 11

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TABLES	» HUMAN	HOMOFOGALE VI	NOTATIONS		Atty. Docket No. 44921-5089WC Doc. No. 1798897.1
Seq. (D) No.	ldenlifer	ConBank Ace./ Ref. Seq. ID No.	Model Gode	Homologous Cane : # Name	Homologous Cluster Name
1633	20713	NM_016999	t	cytochrome P450, subfamily IV B, polypeptide 1, cytochrome P450, subfamily IVB, polypeptide 1	Mus musculus, Similar to cytochrome P450, 4a10, clone MGC:25972 IMAGE:4240359, mRNA, complete cds, RIKEN cDNA A230105L22 gene, cytochrome P450, 4a10, cytochrome P450, 4a14, cytochrome P450, subfamily IVA, polypeptide 11
1633	20711	NM_016999	q,t	1, cytochrome P450,	Mus musculus, Similar to cytochrome P450, 4a10, clone MGC:25972 IMAGE:4240359, mRNA, complete cds, RIKEN cDNA A230105L22 gene, cytochrome P450, 4a10, cytochrome P450, 4a14, cytochrome P450, subfamily IVA, polypeptide 11
1633	20715	NM 016999	q,t	1, cytochrome P450,	Mus musculus, Similar to cytochrome P450, 4a10, clone MGC:25972 IMAGE:4240359, mRNA, complete cds, RIKEN cDNA A230105L22 gene, cytochrome P450, 4a14, cytochrome P450, subfamily IVA, polypeptide 11
1634	1698	NM_017000	e,n,p,General	diaphorase (NADH/NADPH) (cytochrome b-5 reductase), diaphorase 4 (NADH/NADPH)	ESTs, Weakly similar to DHQU RAT NAD(P)H DEHYDROGENASE [R.norvegicus], NAD(P)H menadione oxidoreductase 2, dioxin inducible, NAD(P)H menadione oxidoreductase 2, dioxin-inducible, diaphorase (NADH/NADPH) (cytochrome b-5 reductase), diaphorase 4 (NADH/NADPH)
1635	1399	NM 017006	h,n,General		glucose-6-phosphate dehydrogenase, glucose-6-phosphate dehydrogenase 2, glucose-6-phosphate dehydrogenase X-linked, hexose-6- phosphate dehydrogenase (glucose 1- dehydrogenase)
1000	1000	1411_017000	ii,ii,Oeliciai	glutathione S-transferase A2, glutathione S-	glutathione S-transferase A2, glutathione S-transferase, alpha 2
1637	18989	NM_017013	n	glutathione S-transferase	(Yc2) ESTs, Moderately similar to GLUTATHIONE S-TRANSFERASE YB1 [R.norvegicus], glutathione S- transferase M1, glutathione S-
1638	21013	NM_017014	e,f	transferase, mu 2 glutathione S-transferase M2 (muscle), glutathione S-	transferase, mu 1 ESTs, Moderately similar to GLUTATHIONE S-TRANSFERASE YB1 [R.norvegicus], glutathione S- transferase M1, glutathione S-
1639	11836	NM_017014	e,General	potassium inwardly-	transferase, mu 1 EST, Weakly similar to IRKF MOUSE ATP-SENSITIVE INWARD RECTIFIER POTASSIUM CHANNEL 15 [M.musculus], potassium inwardly- rectifying channel, subfamily J, member 1, potassium inwardly- rectifying channel, subfamily J, member 10, potassium inwardly- rectifying channel, subfamily J, member 15

TABLE	RAMUMAN	HOWOFOGNE V	NOTATIONS		Atty. Docket No. 44921-5089WC Doc. No. 1798897.1
Seq. (ID) No.	ldentiter	Cenbank Ace. Ref. Seq. 10 No.	Model Code.	Honologous Come Name	
1639	5475	NM_017023	b	potassium inwardly- rectifying channel, subfamily J, member 1	EST, Weakly similar to IRKF MOUSE ATP-SENSITIVE INWARD RECTIFIER POTASSIUM CHANNEL 15 [M.musculus], potassium inwardly-rectifying channel, subfamily J, member 1, potassium inwardly-rectifying channel, subfamily J, member 10, potassium inwardly-rectifying channel, subfamily J, member 15
1639	25546	NM_017023	b,bb		ESTs, Highly similar to DEHULM L-
1640	17807	NM_017025	i,General	lactate dehydrogenase 1, A chain, lactate dehydrogenase A	lactate dehydrogenase [H.sapiens], ESTs, Moderately similar to DEHULM L-lactate dehydrogenase [H.sapiens], ESTs, Moderately similar to L-LACTATE DEHYDROGENASE M CHAIN [R.norvegicus], ESTs, Weakly similar to DEMSLM L-lactate dehydrogenase [M.musculus], Homo sapiens, Similar to lactate dehydrogenase 1, A chain, clone MGC:23940 IMAGE:3935569, mRNA, complete cds, expressed sequence Al326310, lactate dehydrogenase 1, A chain, lactate dehydrogenase A, lactate dehydrogenase C
10.10	17707	0 02.0	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	auty er egenado r	Mus musculus adult female placenta
1641	24597	NM_017040	u	protein phosphatase 2 (formerly 2A), catalytic subunit, beta isoform, protein phosphatase 2a, catalytic subunit, beta isoform	cDNA, RIKEN full-length enriched library, clone:1600017J22, full insert sequence, RIKEN cDNA 2310003C10 gene, expressed sequence Al115466, protein phosphatase 2 (formerly 2A), catalytic subunit, beta isoform, protein phosphatase 2a, catalytic subunit, beta isoform, protein phosphatase 4, catalytic subunit
1642	24696	NM_017048	f.j.,z	solute carrier family 4 (anion exchanger), member 2, solute carrier family 4, anion exchanger, member 2 (erythrocyte membrane protein band 3- like 1)	ESTs, Moderately similar to A25104 band 3 protein, nonerythroid [H.sapiens], Human DNA sequence from clone RP4-79416 on chromosome 20 Contains a gene for a putative oncogene protein, parts of 2 novel genes, ESTs, STSs, GSSs and CpG islands, solute carrier family 4 (anion exchanger), member 2, solute carrier family 4, anion exchanger, member 2 (erythrocyte membrane protein band 3-like 1), solute carrier family 4, sodium bicarbonate cotransporter, member 4, solute carrier family 4, sodium bicarbonate cotransporter, member 9, solute carrier family 4, sodium bicarbonate transporter-like, member 11

TABLES	RIMAMUH S	HOWOFOGME W	ENOTATIONS :		Atty. Docket No. 4/1921-5039W0 Doc. No. 1793897.1
	(dentifier	ConBink Ace./ Ref. Seq. ID No.	Medel Gode	Homologous Gene Name	Homologovis Gluster Name
		, a contract of the contract o			EST, Highly similar to B3A3 RAT ANION EXCHANGE PROTEIN 3 [R.norvegicus], ESTs, Weakly similar to I38496 anion exchanger 3 brain isoform [H.sapiens], Human DNA
					sequence from clone RP4-794l6 on chromosome 20 Contains a gene for a putative oncogene protein, parts of 2 novel genes, ESTs, STSs, GSSs and CpG islands, solute carrier family 4 (anion exchanger), member 3, solute
1643	24695	NM_017049	u		carrier family 4, anion exchanger, member 3, solute carrier family 4, sodium bicarbonate transporter-like, member 11
				superoxide dismutase 1,	ESTs, Moderately similar to SUPEROXIDE DISMUTASE [M.musculus], copper chaperone for
				soluble, superoxide dismutase 1, soluble (amyotrophic lateral	superoxide dismutase, superoxide dismutase 1, soluble, superoxide dismutase 1, soluble (amyotrophic
1644	20876	NM_017050	j,n,z	sclerosis 1 (adult)) BCL2-associated X protein, Bcl2-associated X	lateral sclerosis 1 (adult)) BCL2-associated X protein, Bcl2-
1645	910	NM_017059	f,l,m	protein BCL2-associated X	associated X protein
1645	912	NM_017059	i	protein, Bcl2-associated X protein	BCL2-associated X protein, Bcl2- associated X protein ESTs, Moderately similar to
					LYOX_HUMAN PROTEIN-LYSINE 6- OXIDASE PRECURSOR [H.sapiens], ESTs, Moderately similar to PROTEIN- LYSINE 6-OXIDASE PRECURSOR [R.norvegicus], hypothetical protein FLJ21889, lysyl oxidase, lysyl oxidase-
1646	1946	NM_017061	h	lysyl oxidase	like, lysyl oxidase-like 1, lysyl oxidase- like 2 ESTs, Moderately similar to
4545	4042	NM 017061	160	hand avideos	LYOX_HUMAN PROTEIN-LYSINE 6-OXIDASE PRECURSOR [H.sapiens], ESTs, Moderately similar to PROTEIN-LYSINE 6-OXIDASE PRECURSOR [R.norvegicus], hypothetical protein FLJ21889, lysyl oxidase-like, lysyl oxidase-like, lysyl oxidase-
1646	1942	U1/061	t,General	lysyl oxidase	like 2 ESTS, Moderately similar to LYOX_HUMAN PROTEIN-LYSINE 6- OXIDASE PRECURSOR [H.sapiens], ESTS, Moderately similar to PROTEIN- LYSINE 6-OXIDASE PRECURSOR [R.norvegicus], hypothetical protein FLJ21889, lysyl oxidase, lysyl oxidase- like, lysyl oxidase-like 1, lysyl oxidase-
1646	1943	NM_017061	t	lysyl oxidase	like 2 EST, Moderately similar to JH0385 midkine precursor [H.sapiens], EST, Weakly similar to PTN MOUSE PLEIOTROPHIN PRECURSOR [R.norvegicus], midkine, midkine
1647	6062	NM_017066	d	pleiotrophin, pleiotrophin (heparin binding growth factor 8, neurite growth- promoting factor 1)	(neurite growth-promoting factor 2), pleiotrophin, pleiotrophin (heparin binding growth factor 8, neurite growth- promoting factor 1)

TABLES	RIPMENT .	HOWOTOGALS W	ROTATIONS		Miy. Docket No. 44921-5039WG Doc. No. 1793397.1
539. [D] No.	ldentifer	GenBenk Ace./ Ref. Seq. (D No.)	Model Code	Homologous Cene Neme	Homologous Cluster Namo
				lysosomal membrane	CD68 antigen, ESTs, Highly similar to LMP2 RAT LYSOSOME-ASSOCIATED MEMBRANE GLYCOPROTEIN 2 PRECURSOR [R.norvegicus], ESTs, Weakly similar to A48042 lysosomal membrane glycoprotein lamp-2 homolog
1648	6654	NM_017068	w	glycoprotein 2, lysosomal- associated membrane protein 2	[H.sapiens], lysosomal membrane glycoprotein 2, lysosomal-associated membrane protein 2
1649	11153	NM_017073	s	glutamate-ammonia ligase (glutamine synthase), glutamine synthetase	glutamate-ammonia ligase (glutamine synthase), glutamine synthetase
1650	923	NM 017076	General		DNA segment, Chr 7, ERATO Doi 458, expressed, RIKEN cDNA 2610301B19 gene, RIKEN cDNA 3830421F03 gene, poliovirus receptor, poliovirus receptor-related 2 (herpesvirus entry mediator B), poliovirus receptor-related 3, poliovirus sensitivity, tumorassociated antigen 1
1651	1523	NM_017079	s		CD1B antigen, b polypeptide, CD1D antigen, d polypeptide, CD1E antigen, e polypeptide, CD1d1 antigen, CD1d2 antigen
				hydroxysteroid (11-beta) dehydrogenase 1, hydroxysteroid 11-beta	DNA segment, Chr 14, University of California at Los Angeles 2, ESTs, Weakly similar to CORTICOSTEROID 11-BETA-DEHYDROGENASE, ISOZYME 1 [R.norvegicus], expressed sequence C79874, hydroxysteroid (11-beta) dehydrogenase 1, hydroxysteroid 11-beta dehydrogenase 1, hydroxysteroid 17-beta dehydrogenase 11, retinal shortchain dehydrogenase/reductase
1652	23660	NM_017080	s	dehydrogenase 1	retSDR2 Mus musculus, Similar to
1653	275	NM_017081	b,d,General	hydroxysteroid (11-beta) dehydrogenase 2, hydroxysteroid 11-beta dehydrogenase 2	hydroxysteroid 11-beta dehydrogenase 2, clone MGC:25647 IMAGE:4235545, mRNA, complete cds, hydroxysteroid (11-beta) dehydrogenase 2
				uromodulin, uromodulin	RIKEN cDNA 2310037I18 gene, tectorin beta, uromodulin, uromodulin (uromucoid, Tamm-Horsfall glycoprotein), zona pellucida
1654	16211	NM_017082	j,s,z	glycoprotein) glycine N-	glycoprotein 1
1655	1552	NM_017084	<u>j</u>	methyltransferase	glycine N-methyltransferase
1655	1550	NM_017084	у	glycine N- methyltransferase	glycine N-methyltransferase
1656	22552	NM_017087	a,k,x	biglycan	ESTs, Weakly similar to BONE/CARTILAGE PROTEOGLYCAN I PRECURSOR [R.norvegicus], ESTs, Weakly similar to PGS1_HUMAN BONE/CARTILAGE PROTEOGLYCAN I PRECURSOR [H.sapiens], asporin, asporin (LRR class 1), biglycan, opticin

TABLES	HUMAN	HOWO FOCATE (V)	SKIOTIATIONS		Aiiy. Docket No. 44921-5039WO Doc. No. 1793397.1
Seq. ID No.	ldentifier	CenBenk Acc./ Ref. Seq. ID No.	Model Gode	Honologous Cene Name ¥	Homologous Cluster Name
	last resident reconstructors and a	Egolistica in in accomming Amiliano (1992). See See Hills.		The company of the co	ESTs, Weakly similar to ANPA MOUSE ATRIAL NATRIURETIC
					PEPTIDE RECEPTOR A PRECURSOR [M.musculus], ESTs, Weakly similar to GUANYLATE
					CYCLASE SOLUBLE, ALPHA-1
				•	CHAIN [R.norvegicus], guanylate cyclase 1, soluble, alpha 2, guanylate
					cyclase 1, soluble, alpha 3, guanylate cyclase 1, soluble, beta 3, natriuretic
1657	8888	NM 017090	m	guanylate cyclase 1, soluble, alpha 3	peptide receptor 1, soluble guanylyl cyclase alpha2 subunit
1658	10887	NM 017094	a,General	growth hormone receptor	growth hormone receptor
1000	10001		4,000,01	g	ESTs, Highly similar to A Chain A, Cyclophilin A [H.sapiens], ESTs, Weakly similar to PEPTIDYL-PROLYL
					CIS-TRANS ISOMERASE A [R.norvegicus], KIAA1228 protein, RIKEN cDNA 2310076N22 gene, RIKEN cDNA 4930520F12 gene,
					expressed sequence Al256741, expressed sequence AW457192,
				peptidylprolyl isomerase A,	peptidylprolyl isomerase A, peptidylprolyl isomerase A (cyclophilin
1659	4393	NM 017101	a,y		A), peptidylprolyl isomerase E (cyclophilin E)
1009	4000	14W_017 101	a,y	(Cycloprillin A)	blood-brain barrier specific anion
					transporter, solute carrier family
					(organic anion transporter) member
	ļ				10, solute carrier family 21 (organic anion transporter), member 1, solute
					carrier family 21 (organic anion
					transporter), member 10, solute carrier
	ļ				family 21 (organic anion transporter),
					member 14, solute carrier family 21
	1				(organic anion transporter), member 3, solute carrier family 21 (organic anion
			1	solute carrier family 21	transporter), member 6, solute carrier
		// o			family 21 (organic anion transporter),
1660	24770	NM_017111	d	member 1	member 8
1661	20745	NM_017113	е	granulin	granulin
1661	20746	NM_017113	а	granulin	granulin
					DNA segment, Chr 15, ERATO Doi 412, expressed, ESTs, Highly similar to HIPP RAT NEURON SPECIFIC
					CALCIUM-BINDING PROTEIN HIPPOCALCIN [R.norvegicus], ESTs,
					Highly similar to VIS3 MOUSE VISININ LIKE PROTEIN 3 [M.musculus],
					hippocalcin, hippocalcin-like 1,
1662	1375	NM_017122	w	hippocalcin	hypothetical protein FLJ20481
					CD37 antigen, EST, Highly similar to A47629 cell surface glycoprotein CD37
				*	[H.sapiens], Mus musculus, Similar to CD37 antigen, clone MGC:7983
1662	12903	NM_017124	k	CD37 antigen	IMAGE:3585492, mRNA, complete cds, oculospanin
1663	12303	114W V 17 124	II.	Jour anagen	ous, oculospariin

VABLES) HUMAN	HOMOLOGUEAN	SKOLKALOKI	A control of the cont	Atty, Docket No. 44921-5039WO Doc. No. 1793997.1
Seq. (D No.	ldentifer	ConBank Acel Ref. Seq. ID No.	Model Gode	Honologous Cene Neme	Homologous Gluster Name
					EST, Moderately similar to 40S
					RIBOSOMAL PROTEIN SA [R.norvegicus], ESTs, Highly similar to
1					A31233 ribosomal protein RS.40K,
	1				cytosolic [H.sapiens], ESTs, Highly
					similar to A56880 laminin receptor,
					67K [H.sapiens], ESTs, Moderately similar to A29395 ribosomal protein
1					RS.40K - mouse [M.musculus], ESTs,
					Weakly similar to 1405340A protein
					40kD [M.musculus], Homo sapiens
					laminin receptor-like protein LAMRL5
1664	24885	NM 017138	ļ.	laminin receptor 1 (67kD, ribosomal protein SA)	mRNA, complete cds, laminin receptor 1 (67kD, ribosomal protein SA)
1004	24663	14141_017136		Industrial protein SA)	EST, Moderately similar to 40S
					RIBOSOMAL PROTEIN SA
					[R.norvegicus], ESTs, Highly similar to
					A31233 ribosomal protein RS.40K,
					cytosolic [H.sapiens], ESTs, Highly similar to A56880 laminin receptor,
					67K [H.sapiens], ESTs, Moderately
					similar to A29395 ribosomal protein
				r	RS.40K - mouse [M.musculus], ESTs,
					Weakly similar to 1405340A protein
					40kD [M.musculus], Homo sapiens laminin receptor-like protein LAMRL5
				laminin receptor 1 (67kD,	mRNA, complete cds, laminin receptor
1664	24886	NM 017138	d,q	ribosomal protein SA)	1 (67kD, ribosomal protein SA)
					EST, Moderately similar to
					COF1_HUMAN COFILIN, NON-
					MUSCLE ISOFOR [H.sapiens], ESTs,
		İ			Highly similar to DEST_HUMAN DESTRIN [H.sapiens], ESTs,
					Moderately similar to COF1 RAT
					COFILIN, NON-MUSCLE ISOFORM
					[R.norvegicus], ESTs, Moderately
					similar to COF1_HUMAN COFILIN, NON-MUSCLE ISOFOR [H.sapiens],
					cofilin 1 (non-muscle), cofilin 1, non-
				cofilin 1 (non-muscle),	muscle, cofilin 2 (muscle), cofilin 2,
1665	15363	NM_017147	n,u	cofilin 1, non-muscle	muscle, destrin
					EST, Moderately similar to CYSR RAT
					CYSTEINE-RICH PROTEIN 1 [R.norvegicus], ESTs, Weakly similar
					to CYSR RAT CYSTEINE-RICH
					PROTEIN 1 [R.norvegicus], ESTs,
					Weakly similar to S12658 cysteine-rich
					protein [H.sapiens], cysteine and
					glycine-rich protein 1, cysteine rich protein, cysteine-rich protein 2,
					cysteine-rich protein 3, epithelial
					protein lost in neoplasm beta, thymus
1666	13392	NM_017148	u,General		LIM protein

TABLE	E HUMAN	HOMOLOGUE AN	NOTATIONS		Airy, Docker No. 44921-51139/WO Doc. No. 1793597.1
Seg. ID No.	ldeniliter.	GenBenk Acel Ref. Seq. 10 No.	Model Gode	Hemologous Cene Neme	Homologous Cluster Name
					EST, Moderately similar to 60S RIBOSOMAL PROTEIN L29 [R.norvegicus], EST, Weakly similar to S65784 ribosomal protein L29, cytosolic [H.sapiens], ESTs, Highly
					similar to S65784 ribosomal protein L29, cytosolic [H.sapiens], ESTs, Weakly similar to 60S RIBOSOMAL PROTEIN L29 [M.musculus], Human DNA sequence from clone RP4-
					595K12 on chromosome 1p31.2-31.3 Contains a pseudogene similar to 60S RPL29 (ribosomal protein L29 (cell surface heparin binding protein HIP)), a chromosome 1 specific mRNA
1667	5351	NM_017150	q	ribosomal protein L29	(KIAA0499), a novel mRNA (KIAA0433), ESTs, STSs, GSSs and a CpG Island, ribosomal protein L29
					EST, Moderately similar to R3HU15 ribosomal protein S15, cytosolic [H.sapiens], EST, Weakly similar to 40S RIBOSOMAL PROTEIN S15 [R.norvegicus], ESTs, Highly similar to R3HU15 ribosomal protein S15, cytosolic [H.sapiens], Homo sapiens,
1668	16954	NM_017151	a,n	ribosomal protein S15	clone IMAGE:4479080, mRNA, partial cds, ribosomal protein S15 EST, Weakly similar to 40S
					RIBOSOMAL PROTEIN S17 [M.musculus], ESTs, Highly similar to R4HU17 ribosomal protein S17, cytosolic [H.sapiens], ESTs, Moderately similar to R4HU17
1669	21643	NM_017152	g	ribosomal protein S17	ribosomal protein S17, cytosolic [H.sapiens], ribosomal protein S17 EST, Weakly similar to RS3A MOUSE
				ribosomal protein S3A,	40S RIBOSOMAL PROTEIN S3A [M.musculus], ESTs, Highly similar to JC4662 ribosomal protein S3a, cytosolic [H.sapiens], ribosomal
1670	1694	NM_017153	a,q	ribosomal protein S3a	protein S3A, ribosomal protein S3a EST, Moderately similar to R3HU6
					ribosomal protein S6, cytosolic [H.sapiens], EST, Weakly similar to 40S RIBOSOMAL PROTEIN S6 [R.norvegicus], ESTs, Moderately similar to 40S RIBOSOMAL PROTEIN S6 [M.musculus], Homo sapiens cDNA: FLJ23534 fis, clone LNG06974, highly similar to HUMRPS6A Human ribosomal protein S6 mRNA, RIKEN cDNA 5830405M20 gene, ribosomal
1671	17104	NM_017160	bb,General	ribosomal protein S6	protein S6 EST, Moderately similar to R3HU6
					ribosomal protein S6, cytosolic [H.sapiens], EST, Weakly similar to 40S RIBOSOMAL PROTEIN S6 [R.norvegicus], ESTs, Moderately similar to 40S RIBOSOMAL PROTEIN S6 [M.musculus], Homo sapiens cDNA: FLJ23534 fis, clone LNG06974, highly similar to HUMRPS6A Human ribosomal protein S6 mRNA, RIKEN cDNA 5830405M20 gene, ribosomal
1671	17106	NM_017160	u	ribosomal protein S6	protein S6

TABLE	: HUMAN	HOMOLOGUEAK	ROTATIONS		Atty. Docket No. 4/1921-5039WO Doc. No. 1793397.1
Seq. (D. No.		Centenk (Acc./). Ref. Seq. ID No.	Model Gode	Homologove Gane Memo	Homologous Auster Name
					EST, Moderately similar to R3HU6 ribosomal protein S6, cytosolic [H.sapiens], EST, Weakly similar to 40S RIBOSOMAL PROTEIN S6 [R.norvegicus], ESTs, Moderately similar to 40S RIBOSOMAL PROTEIN S6 [M.musculus], Homo sapiens cDNA: FLJ23534 fis, clone LNG06974, highly similar to HUMRPS6A Human ribosomal protein S6 mRNA, RIKEN
1671	17107	NM_017160	d,e	ribosomal protein S6	cDNA 5830405M20 gene, ribosomal protein S6
				glutathione peroxidase 4, glutathione peroxidase 4 (phospholipid	EST, Moderately similar to T02747 phospholipid-hydroperoxide glutathione peroxidase [H.sapiens], EST, Weakly similar to T02747 phospholipid-hydroperoxide glutathione peroxidase [H.sapiens], ESTs, Weakly similar to GSHH RAT PHOSPHOLIPID HYDROPEROXIDE GLUTATHIONE PEROXIDASE [R.norvegicus], Homo sapiens PRO2893 mRNA, complete cds, RIKEN cDNA 2310016C16 gene, RIKEN cDNA 3110050F08 gene, glutathione peroxidase 4, glutathione peroxidase 4 (phospholipid
1672	17686	NM_017165	n,q	(phospholipid hydroperoxidase)	hydroperoxidase)
1673	20702	NM 017166	С	leukemia-associated gene, stathmin 1/oncoprotein 18	ESTs, Weakly similar to STHM MOUSE STATHMIN [M.musculus], Homo sapiens (clone B3B3E13) Huntington's disease candidate region mRNA fragment, leukemia-associated gene, stathmin 1/oncoprotein 18
1674	3513	NM_017177	Г	choline kinase-like, choline/ethanolamine kinase	EST, Weakly similar to KICE MOUSE CHOLINE/ETHANOLAMINE KINASE [M.musculus], ESTs, Weakly similar to KICE RAT CHOLINE/ETHANOLAMINE KINASE [R.norvegicus], Homo sapiens, Similar to hypothetical protein FLJ10761, clone MGC:19512 IMAGE:4329734, mRNA, complete cds, Mus musculus mRNA for choline/ethanolamine kinase, complete cds, RIKEN cDNA 4930555L11 gene, choline kinase-like, choline/ethanolamine kinase, ethanolamine kinase, ethanolamine kinase, hypothetical protein FLJ10761
1675	19031	NM_017180	v,General	T-cell death associated gene, pleckstrin homology- like domain, family A, member 1	ESTs, Weakly similar to S58222 PQrich protein [H.sapiens], MARCKS-like protein, Mus musculus 8 days embryo cDNA, RIKEN full-length enriched library, clone:5730519L10, full insert sequence, Myristoylated alanine-rich protein kinase C substrate, myristoylated alanine rich protein kinase C substrate, myristoylated alanine-rich protein kinase C substrate (MARCKS, 80K-L), pleckstrin homology-like domain, family A, member 1, pleckstrin homology-like domain, family A, member 3, tumor suppressing subtransferable candidate 3

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TABLES	S KAMMAKI S	HOMOLOGUEAN	NOTATIONS		Aity, Docket No. 44921-5089W0 Doc. No. 1798397.1
Seg. ID No.	ldentiliter	GenBank Acc./ Ref. Seq. 10 No.	Model Gode	Homologous Gene Neme	Homologous Gluster Namo
					EST, Moderately similar to HMG2 RAT HIGH MOBILITY GROUP PROTEIN HMG2 [R.norvegicus], EST, Weakly similar to HMG2 RAT HIGH MOBILITY GROUP PROTEIN HMG2 [R.norvegicus], ESTs, Weakly similar to HMG2 RAT HIGH MOBILITY GROUP PROTEIN HMG2 [R.norvegicus], ESTs, Weakly similar to HMG2 RAT HIGH MOBILITY GROUP PROTEIN HMG2 [R.norvegicus], Human DNA sequence from clone RP3-527B10 on chromosome 6q25.1-25.3 Contains a pseudogene similar to HMG (high mobility group) protein, STSs and GSSs, Human DNA sequence from clone RP5-1007G16 on chromosome 1p34.2-35.3. Contains part of the gene for a novel CUB and Sushi (SCR repeat) domain protein, a novel high-mobility group (nonhistone chromosomal) protein 2 (HMG2) like protein (pseudo) gene, a heat shock 60kD protein 1 (chaperonin) (HSPD1) pseudogene, ESTs, STSs and GSSs, RIKEN cDNA 2610021J01 gene, expressed sequence Al326135, expressed sequence Al326135, expressed sequence Al326139, high
					expressed sequence C80539, high mobility group box 2, high-mobility group (nonhistone chromosomal)
1676	15437	NM_017187	x,z		protein 2 EST, Moderately similar to HMG2 RAT HIGH MOBILITY GROUP PROTEIN HMG2 [R.norvegicus], EST, Weakly similar to HMG2 RAT HIGH MOBILITY GROUP PROTEIN HMG2 [R.norvegicus], ESTs, Weakly similar to HMG2 RAT HIGH MOBILITY GROUP PROTEIN HMG2 [R.norvegicus], ESTs, Weakly similar to 201363A high mobility group protein 2 [H.sapiens], ESTs, Weakly similar to HMG2 RAT HIGH MOBILITY GROUP PROTEIN HMG2 [R.norvegicus], Human DNA sequence from clone RP3-527B10 on chromosome 6q25.1-25.3 Contains a pseudogene similar to HMG (high mobility group) protein, STSs and GSSs, Human DNA sequence from clone RP5-1007G16 on chromosome 1p34.2-35.3. Contains part of the gene for a novel CUB and Sushi (SCR repeat) domain protein, a novel highmobility group (nonhistone chromosomal) protein 2 (HMG2) like protein (pseudo) gene, a heat shock 60kD protein 1 (chaperonin) (HSPD1) pseudogene, ESTs, STSs and GSSs, RIKEN cDNA 2610021J01 gene, expressed sequence Al326135, expressed sequence C80539, high mobility group box 2, high-mobility
1676	15433	NM_017187	y		group (nonhistone chromosomal) protein 2

TABLE 3	i Humani	HOMOLOGUEAK	NOTATIONS	TENNY AT 19	Affy: Docket No. 44921-5039W0 Doc. No. 1793397.1
809. ID No.	ldentifier	Genbank Acc./ Ref. Seq. (D No.	Model Code	Homologous Cene Name :	Homologous Gluster Name
					EST, Moderately similar to HMG2 RAT HIGH MOBILITY GROUP PROTEIN HMG2 [R.norvegicus], EST, Weakly similar to HMG2 RAT HIGH MOBILITY GROUP PROTEIN HMG2 [R.norvegicus], ESTs, Weakly similar to 2001363A high mobility group protein 2 [H.sapiens], ESTs, Weakly similar to HMG2 RAT HIGH MOBILITY GROUP PROTEIN HMG2 [R.norvegicus], Human DNA sequence from clone RP3-527810 on chromosome 6q25.1-25.3 Contains a pseudogene similar to HMG (high mobility group) protein, STSs and GSSs, Human DNA sequence from clone RP5-1007G16 on chromosome 1p34.2-35.3. Contains part of the gene for a novel CUB and Sushi (SCR repeat) domain protein, a novel high-mobility group (nonhistone chromosomal) protein 2 (HMG2) like protein (pseudo) gene, a heat shock 60kD protein 1 (chaperonin) (HSPD1) pseudogene, ESTs, STSs and GSSs, RIKEN cDNA 2610021J01 gene, expressed sequence Al326135, expressed sequence C80539, high mobility group box 2, high-mobility group (nonhistone chromosomal)
1676	15434	NM_017187	x,z		protein 2 CD33 antigen, CD33 antigen (gp67),
				malignancy-associated protein, myelin-associated	ESTs, Weakly similar to MYELIN- ASSOCIATED GLYCOPROTEIN PRECURSOR [R.norvegicus], Homo sapiens HSPC078 mRNA, partial cds, myelin associated glycoprotein, myelin-
1677	24437	NM_017190	P	L-kynurenine/alpha- aminoadipate aminotransferase, kynurenine	associated glycoprotein ESTs, Weakly similar to S48737 kynurenine aminotransferase - rat [R.norvegicus], cysteine conjugate- beta lyase; cytoplasmic (glutamine transaminase K, kyneurenine aminotransferase), hypothetical protein
1678	1542	NM_017193	j,i,m,z	aminotransferase II	EST, Weakly similar to COX4_HUMAN CYTOCHROME C OXIDASE POLYPEPTIDE IV PRECURSO [H.sapiens], cytochrome c oxidase subunit IV, cytochrome c oxidase subunit IV isoform 2, cytochrome c oxidase subunit IV isoform 2
1679	14695	NM_017202	q,s		precursor, cytochrome c oxidase, subunit IVa, cytochrome c oxidase, subunit IVb, expressed sequence AL024441 EST, Weakly similar to COX4_HUMAN CYTOCHROME C OXIDASE
1679	14694	NM_017202	s,z		POLYPEPTIDE IV PRECURSO [H.sapiens], cytochrome c oxidase subunit IV, cytochrome c oxidase subunit IV isoform 2, cytochrome c oxidase subunit IV isoform 2 precursor, cytochrome c oxidase, subunit IVa, cytochrome c oxidase, subunit IVb, expressed sequence AL024441

TABLE 3	HUMAN	HOMOLOGUEAX	SKOTKATIONS		Atty., Docket No. 44921-5039WO
Seq. ID	/** 14 5	Conbank Ace/		Homologous Gans	Doc. No. 179339 <u>7.1</u>
No.	Identifier	Ref. Seq. ID No.	Model Code	Name: 11	Homologous Cluster Name ESTs, Highly similar to T09400 outer
					dense fiber protein 2 - mouse
7 0 7					[M.musculus], KIAA1229 protein,
					Myosin heavy chain 11, Myosin, heavy polypeptide 9, non-muscle, expressed
					sequence C80049, myosin, heavy
				outer dense fiber of sperm	polypeptide 9, non-muscle, outer
1600	4400	NIM 017212	m	tails 2, outer dense fibre of sperm tails 2	dense fiber of sperm tails 2, outer dense fibre of sperm tails 2
1680	1428	NM_017213		sperificans 2	ESTs, Moderately similar to 1914205A
					AA transporter [H.sapiens], putative L-
1				a aluda mandan familia 2	type neutral amino acid transporter, solute carrier family 3 (activators of
				solute carrier family 3 (cystine, dibasic and	dibasic and neutral amino acid
				neutral amino acid	transport), member 2, solute carrier
				transporters, activator of	family 3 (cystine, dibasic and neutral
				cystine, dibasic and neutral amino acid transport),	amino acid transporters, activator of cystine, dibasic and neutral amino acid
				member 1, solute carrier	transport), member 1, solute carrier
1681	1622	NM_017216	g,j,s,z	family 3, member 1	family 3, member 1
1682 1682	13642 19976	NM_017220 NM_017220	v w	<u> </u>	
1002	13370	14101-017220			EST, Moderately similar to JC4884
					organic cation transporter protein 2 -
					rat [R.norvegicus], EST, Weakly similar to OCN2 MOUSE ORGANIC
					CATION/CARNITINE TRANSPORTER
					2 [M.musculus], ESTs, Highly similar
					to OCN2_HUMAN ORGANIC CATION/CARNITINE TRANSPORTER
					2 [H.sapiens], ESTs, Highly similar to
					organic cation transporter [H.sapiens],
					solute carrier family 22 (organic cation
1		Ĭ			transporter), member 1, solute carrier family 22 (organic cation transporter),
1					member 4, solute carrier family 22
					(organic cation transporter), member
					5, solute carrier family 22 (organic cation transporter), member 9, solute
					carrier family 22, member 2, solute
1683	1510	NM_017224	General		carrier family 22, member 3
					ESTs, Moderately similar to DRPL RAT ATROPHIN-1 [R.norvegicus],
					ESTs, Weakly similar to G01763
					atrophin-1 [H.sapiens], Homo sapiens,
					clone IMAGE:4153246, mRNA, partial
					cds, RIKEN cDNA 2310009E07 gene, RIKEN cDNA 2810012K09 gene,
				dentatorubral	arginine-glutamic acid dipeptide (RE)
				pallidoluysian atrophy,	repeats, dentatorubral pallidoluysian
		1		dentatorubral- pallidoluysian atrophy	atrophy, dentatorubral-pallidoluysian atrophy (atrophin-1), expressed
1684	1811	NM 017228	j,I,m,z	(atrophin-1)	sequence C78339, formin
					EST, Weakly similar to EFHU2
					translation elongation factor eEF-2
					[H.sapiens], ESTs, Highly similar to ELONGATION FACTOR 2
					[R.norvegicus], ESTs, Weakly similar
					to ELONGATION FACTOR 2
					[R.norvegicus], G1 to phase transition 1, G1 to phase transition 2, RIKEN
					cDNA 4930594C11 gene, U5 snRNP-
					specific protein, 116 kD, eukaryotic
					translation elongation factor 1 alpha 1, eukaryotic translation elongation factor
				eukaryotic translation	2, expressed sequence Al451340,
1686	17563	NM_017245	a,c,e,q	elongation factor 2	hypothetical protein FLJ21661

TABLES		HOMOFOGREYK	SAOITATON		Any. Docket No. 44921-5039W0 Doc. No. 1793397.1
Sଙ୍କୁ, ID No.	loemilier	CenBank Acc.//. Ref. Seq. ID No.!		Homologous Ceita	Homologous Guster Name
					ESTs, Highly similar to ROA1 RAT HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEIN A1 [R.norvegicus], ESTs, Moderately similar to Up1, The Two Rna-Recognition Motif Domain Of Hnrnp A1 (SUB 3-184 [H.sapiens], ESTs, Weakly similar to ROA1 RAT HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEIN A1 [R.norvegicus], ESTs, Weakly similar to heterogeneous ribonuclear particle protein A1 [H.sapiens], Human DNA sequence from clone RP11-51N22 on chromosome 13 Contains ESTs, STSs
.1687	17502	NM 017248	ı		and GSSs. Contains an HNRPA1 (heterogeneous nuclear ribonucleoprotein A1) pseudogene, RIKEN cDNA 4930547K05 gene, heterogeneous nuclear ribonucleoprotein A1
1687	17501	NM_017248	x	heterogeneous nuclear ribonucleoprotein A1 B-cell translocation gene 1,	ESTs, Highly similar to ROA1 RAT HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEIN A1 [R.norvegicus], ESTs, Moderately similar to Up1, The Two Rna-Recognition Motif Domain Of Hnrnp A1 {SUB 3-184 [H.sapiens], ESTs, Weakly similar to ROA1 RAT HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEIN A1 [R.norvegicus], ESTs, Weakly similar to heterogeneous ribonuclear particle protein A1 [H.sapiens], Human DNA sequence from clone RP11-51N22 on chromosome 13 Contains ESTs, STSs and GSSs. Contains an HNRPA1 (heterogeneous nuclear ribonucleoprotein A1) pseudogene, RIKEN cDNA 4930547K05 gene, heterogeneous nuclear ribonucleoprotein A1 B-cell translocation gene 1, antiproliferative, ESTs, Weakly similar to BTG1 RAT BTG1 PROTEIN [R.norvegicus], transducer of ERBB2, 1, transducer
1688	19	NM_017258	v,General	anti-proliferative	of ErbB-2.1 B-cell translocation gene 2, anti- proliferative, B-cell translocation gene
1689	15300	NM_017259	i,v,cc,General	B-cell translocation gene 2, anti-proliferative, BTG family, member 2	3, B-cell translocation gene 4, BTG family, member 2, BTG family, member 3, BTG family, member 4, ESTs, Highly similar to BTG2_HUMAN BTG2 PROTEIN PRECURSOR [H.sapiens] B-cell translocation gene 2, antiproliferative, B-cell translocation gene
1689	15301	NM_017259	I,m,v,aa,cc,Gener al	B-cell translocation gene 2, anti-proliferative, BTG family, member 2	3, B-cell translocation gene 4, BTG family, member 2, BTG family, member 3, BTG family, member 4, ESTs, Highly similar to BTG2_HUMAN BTG2 PROTEIN PRECURSOR [H.sapiens]

TABLES	HUMAN	HONOFOGRE W			Aiiy. Docket No. 44921-5039WO Doc. No. 1793397.1
Seq. ID No.	Identifier	CenBank Ace. Rol. Seq. ID No.	Model Gode	Homologovs Cene Memo	Homologous Gluster Name
1689	15299	NM 017259	l,y,cc,General	B-cell translocation gene 2, anti-proliferative, BTG family, member 2	B-cell translocation gene 2, anti- proliferative, B-cell translocation gene 3, B-cell translocation gene 4, BTG family, member 2, BTG family, member 3, BTG family, member 4, ESTs, Highly similar to BTG2_HUMAN BTG2_PROTEIN PRECURSOR [H.sapiens]
				protease (prosome, macropain) 28 subunit, alpha, proteasome (prosome, macropain) activator subunit 1 (PA28	EST, Moderately similar to A Chain A, Proteasome Activator Reg(Alpha) (SUB 4-63 [H.sapiens], proteasome (prosome, macropain) activator subunit 1 (PA28 alpha)
1691	3987	NM_017264	bb	proteasome (prosome, macropain) subunit, alpha type 3, proteasome (prosome, macropain) subunit, alpha type, 3	multicatalytic endopeptidase complex [H.sapiens], ESTs, Highly similar to PRC8 MOUSE PROTEASOME COMPONENT C8 [M.musculus], ESTs, Highly similar to PROTEASOME COMPONENT C8 [R.norvegicus], ESTs, Weakly similar to SNHUC8 multicatalytic endopeptidase complex [H.sapiens], proteasome (prosome, macropain) subunit, alpha type 3, proteasome (prosome, macropain) subunit, alpha type, 3
				proteasome (prosome, macropain) subunit, alpha type 4, proteasome (prosome, macropain)	EST, Weakly similar to SNHUC9 multicatalytic endopeptidase complex [H.sapiens], proteasome (prosome, macropain) subunit, alpha type 4, proteasome (prosome, macropain)
1692	1447	NM_017281		proteasome (prosome, macropain) subunit, alpha type 6, proteasome (prosome, macropain)	subunit, alpha type, 4 ESTs, Weakly similar to PRCI_HUMAN PROTEASOME IOTA CHAIN [R.norvegicus], proteasome (prosome, macropain) subunit, alpha type 6, proteasome (prosome,
1693	15535	NM_017283	s,bb	subunit, alpha type, 6 ATPase, Ca++ transporting, cardiac	macropain) subunit, alpha type, 6
1694	12349	NM_017290	General	calcium channel, voltage-	EST, Highly similar to CCAD MOUSE VOLTAGE-DEPENDENT L-TYPE CALCIUM CHANNEL ALPHA-1D SUBUNIT [M. musculus], EST, Moderately similar to CCAD MOUSE VOLTAGE-DEPENDENT L-TYPE CALCIUM CHANNEL ALPHA-1D SUBUNIT [M. musculus], EST, Moderately similar to CCAD RAT VOLTAGE-DEPENDENT L-TYPE CALCIUM CHANNEL ALPHA-1D SUBUNIT [R. norvegicus], RIKEN cDNA 8430418G19 gene, calcium channe, voltage-dependent, alpha 1F subunit, calcium channel, voltage-dependent, alpha 1F subunit, calcium channel, voltage-dependent, alpha 1F subunit, rolveystic kidney
1695	15819	NM_017298	p	dependent, L type, alpha 1D subunit	disease 2-like 2

TABLE &	KAMUH 3	HOMOFOGRE VI			Attiy, Docket No. 44921-5089WC Doc. No. 1793397.1
Seg. ID No.	ldentifer	Conean's Aced Ref. Seq. ID No.	Model Code	Homologous Gano Namo	Homologous Cluster Neme
1696	23825	NM 017299	v	solute carrier family 19 (folate transporter), member 1, solute carrier family 19 (sodium/hydrogen exchanger), member 1	expressed sequence AW322295, solute carrier family 19 (folate transporter), member 1, solute carrier family 19 (sodium/hydrogen exchanger), member 1, solute carrier family 19 (sodium/hydrogen exchanger), member 3, solute carrier family 19 (thiamine transporter), member 2, solute carrier family 19, member 3
1696	23826	NM_017299	v	solute carrier family 19 (folate transporter), member 1, solute carrier family 19 (sodium/hydrogen exchanger), member 1	expressed sequence AW322295, solute carrier family 19 (folate transporter), member 1, solute carrier family 19 (sodium/hydrogen exchanger), member 1, solute carrier family 19 (sodium/hydrogen exchanger), member 3, solute carrier family 19 (thiamine transporter), member 2, solute carrier family 19, solute carrier family 19, member 3
1697 1698	14003 26109	NM_017305 NM_017306	j,l,m,y,z lg,s	glutamate-cysteine ligase , modifier subunit, glutamate- cysteine ligase, modifier subunit	ESTs, Highly similar to GSH0_HUMAN GLUTAMATECYSTEINE LIGASE REGULATORY SUBUNIT [H.sapiens], glutamate-cysteine ligase, modifier subunit, glutamate-cysteine ligase, modifier subunit
1698	18687	NM_017306	q,t	dodecenoyl-Coenzyme A delta isomerase (3,2 trans- enoyl-Coenyme A isomerase), dodecenoyl- Coenzyme A delta isomerase (3,2 trans-enoyl- Coenzyme A isomerase)	Coenzyme A isomerase)
1699	18142	NM_017314 NM_017320	g.s,aa		EST, Moderately similar to S12583 polyubiquitin 4 - mouse [M.musculus], EST, Weakly similar to JE0190 polyubiquitin unit [H.sapiens], Homo sapiens UBBP2 pseudogene for ubiquitin UBB, RIKEN CDNA 2700054004 gene, expressed sequence Al194771, expressed sequence AL033289, ubiquitin B, ubiquitin C cathepsin S
1701	20809	NM_017326 NM_017334	u	calmodulin 2, calmodulin 2 (phosphorylase kinase, delta)	Calmodulin 1 (phosphorylase kinase, delta), Calmodulin III, ESTs, Highly similar to A Chain A, Calmodulin Complexed With Calmodulin-Binding Peptide From Smooth Muscle Myosin Light Chain Kinase {SUB 2-148 [H.sapiens], R.norvegicus CaMII retropseudogene (clone lambda SC27), RIKEN cDNA 2310068022 gene, calmodulin 1, calmodulin 1 (phosphorylase kinase, delta), calmodulin 2, calmodulin 2 (phosphorylase kinase, delta), calmodulin 3, calmodulin-like 3

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TABLE 8	KINWWW.	HOMOLOGUEAN	EKIOTEATOKI		Atty, Dockel No. 44921-5039W0 Doc. No. 1793397.1
Seg. [D - No.	ldentifier/	GenBank Ave./ Ref. Seq. ID No.	Model Gode	Homologous Cene Name	Emsk relevið evogolomok
1703	16148	NM_017340	q,s		ESTs, Highly similar to ACYL-COENZYME A OXIDASE, PEROXISOMAL [R.norvegicus], ESTs, Highly similar to CAOP_HUMAN ACYL-COENZYME A OXIDASE, PEROXISOMAL [H.sapiens], RIKEN cDNA 1300003009 gene, RIKEN cDNA 2310016C19 gene, acetyl- Coenzyme A dehydrogenase, short chain, acyl-Coenzyme A oxidase 2, branched chain, isovaleryl coenzyme A dehydrogenase
1703	16150	NM 017340	а		ESTs, Highly similar to ACYL-COENZYME A OXIDASE, PEROXISOMAL [R.norvegicus], ESTs, Highly similar to CAOP_HUMAN ACYL-COENZYME A OXIDASE, PEROXISOMAL [H.sapiens], RIKEN cDNA 1300003009 gene, RIKEN cDNA 2310016C19 gene, acetyl-Coenzyme A dehydrogenase, short chain, acyl-Coenzyme A oxidase 2, branched chain, isovaleryl coenzyme A dehydrogenase
					ESTs, Weakly similar to MOHULP myosin regulatory light chain, placental [H.sapiens], RIKEN cDNA 2900073G15 gene, expressed sequence C77744, myosin, light polypeptide, regulatory, non-
1704	20849	NM_017343	r,u,General		sarcomeric (20kD) ESTs, Weakly similar to MOHULP myosin regulatory light chain, placental [H.sapiens], RIKEN cDNA 2900073G15 gene, expressed sequence C77744, myosin, light polypeptide, regulatory, non-
1704	20848	NM_017343	b,General		sarcomeric (20kD)
					GPI-anchored metastasis-associated protein homolog, metastasis-associated GPI-anchored protein, plasminogen activator, urokinase receptor, urokinase plasminogen
1705	606	NM_017350	b	DD7 and LIM descript	activator receptor ESTs, Weakly similar to LIM PROTEIN CLP36 [R.norvegicus], PDZ and LIM domain 1 (elfin), PDZ and LIM domain 3, PDZ-LIM protein mystique, RIKEN cDNA 1110003B01 gene, Rattus norvegicus LIM-domain protein LMP-1 mRNA, complete cds, Z-band alternatively spliced PDZ-motif, actinin alpha 2 associated LIM protein, alpha-actinin-2-associated LIM protein,
1706	1581	NM_017365	General	PDZ and LIM domain 1 (elfin)	reversion induced LIM gene ESTs, Moderately similar to alpha-
1707	455	NM_019131	x	tropomyosin 1 (alpha), tropomyosin 1, alpha	tropomyosin slow [M.musculus], tropomyosin 4 ESTs, Moderately similar to alpha-
1707	456	NM_019131	y,z	tropomyosin 1 (alpha), tropomyosin 1, alpha	tropomyosin slow [M.musculus], tropomyosin 4

TABLE S	K HJUJMANNI	HOMOLOGUEAN	BUOTATIONS		Anny, Doctet No. 44921-5039W0 Doc. No. 1793397.1
Seq. (D (No.)	(dentifier	Conbook Aced Rof. Seq. ID No.	Model Code	Hemologous Cont Name	Homologovis Cluster Name
(Presidential Presidential				solute carrier family 12 (sodium/potassium/chlorid	EST, Weakly similar to NKC2_HUMAN BUMETANIDE-SENSITIVE SODIUM-(POTASSIUM)-CHLORIDE COTRANSPORTER 2 (KIDNEY-SPECIFIC NA-K-CL SYMPORTER) [H.sapiens], expressed sequence AI788571, hypothetical protein FLJ23188, solute carrier family 12 (sodium/potassium/chloride
1708	4532	NM 019134	Ь	e transporters), member 1, solute carrier family 12, member 1	transporters), member 1, solute carrier family 12, member 1, solute carrier family 12, member 2
					ESTs, Moderately similar to SNG1 RAT SYNAPTOGYRIN 1 [R.norvegicus], synaptogyrin 1,
1709	1608	NM_019166	j,y,z	synaptogyrin 1	synaptogyrin 3, synaptogyrin 4
				synuclein, alpha, synuclein, alpha (non A4 component of amyloid	synuclein, alpha, synuclein, alpha (non A4 component of amyloid precursor),
1710	7489	NM_019169	c,General	precursor)	synuclein, beta, synuclein, gamma ESTs, Weakly similar to JC5284
					carbonyl reductase (NADPH) (EC 1.1.1.184), inducible - rat [R.norvegicus], Homo sapiens, clone MGC:23280 IMAGE:4637504, mRNA, complete cds, RIKEN cDNA 1110001J05 gene, RIKEN cDNA 9430059D04 gene, carbonyl reductase 1, carbonyl reductase 3, expressed
1711	17066	NM_019170	P		sequence C81353 ESTs, Weakly similar to CARBONIC ANHYDRASE IV PRECURSOR [R.norvegicus], carbonic anhydrase 15, carbonic anhydrase 4, carbonic
1712	23924	NM 019174	bb	carbonic anhydrase 4, carbonic anhydrase IV	anhydrase IV, carbonic anhydrase XIV, expressed sequence AW456718
17.12	20027			ADP-ribosylation factor-like	ADP-ribosylation factor-like 4, ADP-ribosylation factor-like 7, ADP-ribosylation-like 4, ESTs, Weakly similar to ARL4 MOUSE ADP-RIBOSYLATION FACTOR-LIKE PROTEIN 4 [M.musculus], Mus musculus, Similar to ADP-ribosylation-
1713	24019	NM_019186	t	4, ADP-ribosylation-like 4	IMAGE:3599701, mRNA, complete cds
1714	22063	NM_019195	d	CD47 antigen (Rh-related antigen, integrin- associated signal transducer), integrin- associated protein	FOT W11
					ESTs, Weakly similar to GRG MOUSE GRG PROTEIN [R.norvegicus], amino
1715	2079	NM_019220	j,k,z		terminal enhancer of split

TABLE 3	: HUMÁN	HOMOTOGNE W	NOTATIONS	Photography and the second sec	Ally, Docket No. 44921-5189WO Doc. No. 1798897.1
Seq. [D No.	Identifier	Consonkace Ref Seq. 10 No.	Model Code	Homologous Gane Name	Homologous Cluster Name
				solute carrier family 12 (potassium/chloride transporters), member 4, solute carrier family 12,	EST, Moderately similar to T31429 K-Cl cotransport protein KCC1, furosemide-sensitive - rat [R.norvegicus], EST, Weakly similar to T31429 K-Cl cotransport protein KCC1, furosemide-sensitive - rat [R.norvegicus], ESTs, Highly similar to T17275 hypothetical protein DKFZp434D2135.1 [H.sapiens], Mus musculus strain ILS K-Cl cotransporter (Slc12a5) mRNA, complete cds, Rattus norvegicus ccc6 mRNA for cation-chloride cotransporter 6, complete cds, expressed sequence AW546649, solute carrier family 12 (potassium/chloride transporters), member 4, solute carrier family 12 (potassium/chloride transporters), member 6, solute carrier family 12 (potassium/chloride transporters), member 7, solute carrier family 12, member 2, solute carrier family 12, member 4, solute carrier family 12, member 4, solute carrier family 12, member 4, solute carrier family 12, member 4, solute carrier family 12,
1716	16284	NM_019229	l,m	member 4 small inducible cytokine	member 7
1717	985	NM 019233	b,cc	subfamily A (Cys-Cys), member 20, small inducible cytokine subfamily A20	small inducible cytokine subfamily A (Cys-Cys), member 20, small inducible cytokine subfamily A20
1718	15503	NM_019237	k,x	procollagen C- endopeptidase enhancer, procollagen C-proteinase enhancer protein	EST, Weakly similar to PCO1_HUMAN PROCOLLAGEN C-PROTEINASE ENHANCER PROTEIN PRECURSOR [H.sapiens], Homo sapiens cDNA FLJ12558 fis, clone NT2RM4000787, Mus musculus CSMD1 (Csmd1) mRNA, complete cds, RIKEN cDNA 2400001018 gene, expressed sequence Al043106, membrane-type frizzled-related protein, procollagen C-endopeptidase enhancer, procollagen C-endopeptidase enhancer 2, procollagen C-protein EST. Weakly similar to PCO1. HUMAN
171P	15504	NM 019237	k,x	procollagen C- endopeptidase enhancer, procollagen C-proteinase enhancer protein	EST, Weakly similar to PCO1_HUMAN PROCOLLAGEN C-PROTEINASE ENHANCER PROTEIN PRECURSOR [H.sapiens], Homo sapiens cDNA FLJ12558 fis, clone NT2RM4000787, Mus musculus CSMD1 (Csmd1) mRNA, complete cds, RIKEN cDNA 2400001018 gene, expressed sequence Al043106, membrane-type frizzled-related protein, procollagen C-endopeptidase enhancer, procollagen C-endopeptidase enhancer 2, procollagen C-proteinase enhancer protein
1718	15504	NM_019237	J.,A	emancer protein	ESTs, Weakly similar to INTERFERON RELATED PROTEIN PC4 [R.norvegicus], interferon-related
1719	17908	NM_019242	l,v,cc,General	interferon-related developmental regulator 1	developmental regulator 1, interferon- related developmental regulator 2
1720	11218	NM_019247	С	paired-like homeodomain transcription factor 3	paired-like homeodomain transcription factor 1, paired-like homeodomain transcription factor 2, paired-like homeodomain transcription factor 3

TABLES	e HUMAN	HOMOLOGUEAN	NOTATIONS		Aity: Docket No. 44921-5039WO Doc No. 1793397.1
Seq. ID No.	nden Miers	GenBank Accul Ref. Seq. ID No.	Model Gode	Homologous Gene Name	Homologous Cluster Name
1721	15259	NM 019259	d,f	Security of the second security of the second secon	DNA segment, Chr 11, Wayne State University 182, expressed, complement component 1, q subcomponent binding protein, expressed sequence AA986492
1721	13239	1411/2013233	<u>u,ı</u>		C1q-related factor, Homo sapiens, Similar to complement component 1, q subcomponent, c polypeptide, clone MGC:17279 IMAGE:4212772, mRNA, complete cds, complement component 1, q subcomponent, beta polypeptide,
1722	21443	NM_019262	aa,General	complement component 1, q subcomponent, beta polypeptide	complement component 1, q subcomponent, c polypeptide, expressed sequence Al385742
1722	21444	NM 019262	t,General	complement component 1, q subcomponent, beta polypeptide	C1q-related factor, Homo sapiens, Similar to complement component 1, q subcomponent, c polypeptide, clone MGC:17279 IMAGE:4212772, mRNA, complete cds, complement component 1, q subcomponent, beta polypeptide, complement component 1, q subcomponent, c polypeptide, expressed sequence Al385742
				sodium channel, voltage gated, type VIII, alpha polypeptide, sodium channel, voltage-gated,	ESTs, Highly similar to voltage gated Na channel Scn8a [M.musculus], Mus musculus adult male hypothalamus cDNA, RIKEN full-length enriched library, clone:A230108N10, full insert sequence, hypothetical protein MGC15619, sodium channel, voltage gated, type VIII, alpha polypeptide, sodium channel, voltage-gated, type II, alpha 2 polypeptide, sodium channel, voltage-gated, type III, alpha polypeptide, sodium channel, voltage-gated, type III, alpha polypeptide, sodium channel, voltage-
1723	117	NM_019266	o,bb	gap junction membrane channel protein alpha 5,	датей, туре VIII, агрна розурершие
1724	1145	NM_019280	w	complex, alcohol	ESTs, Weakly similar to ADHA MOUSE ALCOHOL DEHYDROGENASE A CHAIN [M.musculus], alcohol dehydrogenase 1A (class I), alpha polypeptide, alcohol dehydrogenase 1B (class I), beta polypeptide, expressed sequence
1725	22220	NM_019286	С	dehydrogenase 1A (class I), alpha polypeptide	Al194826, nuclear receptor binding factor 1 EST, Highly similar to AR41_HUMAN ARP2/3 COMPLEX 41 KDA SUBUNIT [H.sapiens], ESTs, Moderately similar to AR41_HUMAN ARP2/3 COMPLEX 41 KDA SUBUNIT [H.sapiens], actin related protein 2/3 complex, subunit 1A (41 kD), actin related protein 2/3 complex, subunit related protein 2/3 complex, subunit 1A (41 kDa), actin related protein 2/3 complex, subunit
1726	10015	NM_019289	l,m,t,x,General		1B (41 kDa), expressed sequence AA408064, suppressor of profilin/p41 of actin-related complex 2/3

TABLES	KAMUHI 3	HOMOLOGAE VIX	SMOTRATIONS		Aity, Docket No. 44921-5039WO Doc. No. 1798397.1
Seq. [D] No.	(den)(liter	Conbonk Ace <i>l</i> Ref. Soq. ID No:	Model Code	Momologous Cana Name	Homologous Cluster Name
		V			EST, Highly similar to AR41_HUMAN ARP2/3 COMPLEX 41 KDA SUBUNIT [H.sapiens], ESTs, Moderately similar to AR41_HUMAN ARP2/3 COMPLEX 41 KDA SUBUNIT [H.sapiens], actin related protein 2/3 complex, subunit 1A (41 kD), actin related protein 2/3 complex, subunit 1A (41 kDa), actin related protein 2/3 complex, subunit 1B (41 kDa), expressed sequence
1726	10016	NM_019289	bb,General		AA408064, suppressor of profilin/p41 of actin-related complex 2/3
					RIKEN cDNA 2310015O17 gene, RIKEN cDNA 4933411O17 gene, cell division cycle 2 homolog A (S. pombe), cell division cycle 2, G1 to S and G2 to M, cyclin-dependent kinase-like 1 (CDC2-related kinase), cyclin- dependent kinase-like 2 (CDC2-related kinase), cyclin-dependent kinase-like 3, expressed sequence Al852479,
1727	21651	NM_019296	c,f,x		serine/threonine kinase NKIATRE beta
1728	20751	NM_019301	s	·	CUB and Sushi multiple domains 1, ESTs, Highly similar to 173012 complement C3b/C4b receptor, membrane-bound form precursor [H.sapiens], ESTs, Weakly similar to JC2054 complement regulatory protein, 5l2 antigen precursor - rat [R.norvegicus], Mus musculus 8 days embryo cDNA, RIKEN full-length enriched library, clone:5730478H20, full insert sequence, complement component (3b/4b) receptor 1, including Knops blood group system, complement receptor related protein, decay accelerating factor 1, decay accelerating factor 2, membrane cofactor protein
1729 1730	645 1301	NM_019345 NM_019349	bb c	solute carrier family 12 (sodium/chloride transporters), member 3, solute carrier family 12, member 3	ESTs, Moderately similar to PC4180 thiazide-sensitive sodium-chloride cotransporter [H.sapiens], ESTs, Moderately similar to thiazide-sensitive Na-Cl cotransporter [M.musculus], ESTs, Weakly similar to BUMETANIDE-SENSITIVE SODIUM-(POTASSIUM)-CHLORIDE COTRANSPORTER 2 [M.musculus], expressed sequence Al788571, solute carrier family 12 (sodium/chloride transporters), member 3, solute carrier family 12, member 3

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TABLE 8	e (Handan	HOMOFOGREYX	- SKOTATOKI		Aity, Docket No. 44921-5039We Doc. No. 1793397.1
Sog. (D No.	ldemiliter	GenBenk Ace <i>l</i> Ref. Seq. (D No.		Homologous Gans Name: 11	Homologous Cluster Name
					ESTs, Moderately similar to BMCP_HUMAN BRAIN MITOCHONDRIAL CARRIER PROTEIN-1 [H.sapiens], RIKEN cDNA 3632410G24 gene, RIKEN cDNA 4933433D23 gene, expressed sequence AW108044, solute carrier family 25 (mitochondrial carrier, brain), member 14, solute carrier family 25 (mitochondrial carrier; adenine nucleotide translocator), member 10, solute carrier family 25 (mitochondrial carrier; ornithine transporter), member 15, uncoupling protein 2 (mitochondrial, proton carrier),
1731	3776	NM_019354	a,u		uncoupling protein 2, mitochondrial RIKEN cDNA 0910001023 gene, eukaryotic translation initiation factor
1732	4592	NM 019356	General		2, subunit 1 (alpha, 35kD), eukaryotic translation initiation factor 2A
1733	1324	NM_019371	w		EGL nine (C.elegans) homolog 1, EGL nine (C.elegans) homolog 2, EGL nine (C.elegans) homolog 3, EGL nine homolog 3 (C. elegans), ESTs, Moderately similar to A53770 growth factor-responsive protein, vascular smooth muscle - rat [R.norvegicus], ESTs, Weakly similar to A53770 growth factor-responsive protein, vascular smooth muscle - rat [R.norvegicus], SCAN domain-containing 2
1734	19577	NM 019377	e	tyrosine 3- monooxgenase/tryptophan 5-monooxgenase activation protein, beta polypeptide, tyrosine 3- monooxygenase/tryptopha n 5-monooxygenase activation protein, beta polypeptide	Containing 2
1735	24626	NM_019381	s	testis enhanced gene transcript, testis enhanced gene transcript (BAX inhibitor 1)	CGI-119 protein, RIKEN cDNA 5031406P05 gene, testis enhanced gene transcript (BAX inhibitor 1)
1736	744	NM 019622	p		ESTs, Highly similar to T42716 ankyrin 3, splice form 4 - mouse [M.musculus], ESTs, Moderately similar to A55575 ankyrin 3, long splice form [H.sapiens], ESTs, Weakly similar to T42716 ankyrin 3, splice form 4 - mouse [M.musculus], RIKEN cDNA 2310026G15 gene, RIKEN cDNA 4833425P12 gene, RIKEN cDNA 4930400E23 gene, RIKEN cDNA C430011H06 gene, ankyrin 3, node of Ranvier (ankyrin G), hypothetical protein FLJ20189, phospholipase A2, group VI (cytosolic, calcium-independent), proteasome (prosome, macropain) 26S subunit, non-ATPase, 10

TABLE		HOMOLOGUE AX			Affy, Docket No. 44921-5039WC Doc. No. 1793397.1
Sog. (D No.	ldentifer	ConBank Acel Ref. Seq. ID No.	Model Gode	Homologous Cene Name	Homologous Gluster Name
				galaite galaite et annual annual annual annual annual annual annual annual annual annual annual annual annual	EST, Weakly similar to CPF1 RAT CYTOCHROME P450 4F1 [R.norvegicus], ESTs, Weakly similar to S45702 leukotriene-B4 20-monooxygenase [H.sapiens], Mus musculus, Similar to RIKEN cDNA 1810054N16 gene, clone MGC:7384 IMAGE:3487830, mRNA, complete cds, RIKEN cDNA 2310021J05 gene, cytochrome P450 isoform 4F12, cytochrome P450, subfamily IV B, polypeptide 1, cytochrome P450, subfamily IVF,
1737	20716	NM_019623	с		polypeptide 2, cytochrome P450, subfamily IVF, polypeptide 8, expressed sequence AI787289 EST. Moderately similar to 1713410A
4770	20700	NIM 040004		lectin, galactose binding, soluble 1, lectin, galactoside-binding, soluble 1 (ralectin 1)	beta galactoside soluble lectin [H.sapiens], EST, Moderately similar to GALECTIN-1 [R.norvegicus], Human HL14 gene encoding beta-galactoside-binding lectin, 3' end, clone 2, RIKEN cDNA 2200008F12 gene, Rattus norvegicus mRNA for galectin-2 related protein, complete cds, lectin, galactose binding, soluble 1, lectin, galactoside-binding, soluble, 1 (galectin 1), lectin, galactoside-binding soluble, 2 (galectin 2)
1739	574	NM_019904	u,General	hydroxyacid oxidase (glycolate oxidase) 3, hydroxyacid oxidase 3 (medium-chain)	ESTS, Highly similar to (S)-2-HYDROXY-ACID OXIDASE, PEROXISOMAL [R.norvegicus], ESTS, Highly similar to LUHU36 annexin II [H.sapiens], RIKEN cDNA 1110003P15 gene, RIKEN cDNA B430311C09 gene, annexin A2, annexin A2 pseudogene 2, caspase recruitment domain family, member 6, expressed sequence AW215814, hydroxyacid oxidase (glycolate oxidase) 3, hydroxyacid oxidase 1, liver, hydroxyacid oxidase 2 (long chain), nucleolar protein 3 (apoptosis repressor with CARD domain)
				hypothetical protein similar to mouse aldehyde reductase 6 (renal), renal-	
1740	9096	NM_019908	j . Canand	specific oxido-reducatse	parathyroid hormone receptor, parathyroid hormone receptor 1
1741	20457	NM_020073 NM 020073	i,General General		parathyroid hormone receptor, parathyroid hormone receptor, parathyroid hormone receptor 1
1741	20458	NM_020073	General		parathyroid hormone receptor, parathyroid hormone receptor,
1742	18713	NM_020075	r	eukaryotic translation initiation factor 5	DNA segment, Chr 12, ERATO Doi 549, expressed, KIAA1856 protein, eukaryotic translation initiation factor 5 DNA segment, Chr 12, ERATO Doi
1742	18715	NM_020075	r	eukaryotic translation initiation factor 5	549, expressed, KIAA1856 protein, eukaryotic translation initiation factor 5 3-hydroxyanthranilate 3,4- dioxygenase, RIKEN cDNA
1743	20493	NM_020076	p		0610007K21 gene, RIKEN cDNA 0610012J07 gene

TABLES	HUMAN	HOMOLOGUEAN	SKOTEATEON	A COMPANY OF A VICE	
					Doc. No. 1798897.1
809. [D No: 7	ldentifier	Consonk Ace./ Ref. Seq. ID No.	Modal (Goda	Homologous Gene * Name	Homologous Cluster Name
					angiotensin I converting enzyme
1744	16275	NIM 020076	_		(peptidyl-dipeptidase A) 2, kidney- specific membrane protein
1744	16375	NM_020976	g		EST, Highly similar to THYMOSIN
					BETA-10 [R.norvegicus], ESTs, Highly
	1				similar to TYB4 MOUSE THYMOSIN
					BETA-4 [M.musculus], expressed
4745	00046	NIM 024264	k Canaral		sequence AW544206, thymosin, beta 10, thymosin, beta 4, X chromosome
1745	20816	NM_021261	k,General		EST, Weakly similar to 60S
					RIBOSOMAL PROTEIN L35A
					[R.norvegicus], EST, Weakly similar to
					R35A MOUSE 60S RIBOSOMAL
					PROTEIN L35A [M.musculus], EST,
					Weakly similar to R5HU35 ribosomal protein L35a [H.sapiens], Homo
					sapiens cDNA FLJ11509 fis, clone
	1				HEMBA1002166, RIKEN cDNA
	l				2810431L15 gene, ribosomal protein
4740	45005	NN 024264		ribosomal protein L35a	L35a, uncharacterized hypothalamus protein HSMNP1
1746	15335	NM_021264	a	transforming growth factor,	protein Howin 1
1747	18729	NM 021578	k,z	beta 1	transforming growth factor, beta 1
					EST, Weakly similar to TGFB_HUMAN
					LATENT TRANSFORMING GROWTH
					FACTOR BETA BINDING PROTEIN 1 PRECURSOR [H.sapiens], ESTs,
	l				Weakly similar to TGFB RAT LATENT
					TRANSFORMING GROWTH FACTOR
					BETA BINDING PROTEIN 1
					PRECURSOR [R.norvegicus], RIKEN
					cDNA 2310046A13 gene, hypothetical protein MGC13010, latent transforming
					growth factor beta binding protein 1,
				latent transforming growth	latent transforming growth factor beta
				factor beta binding protein	binding protein 2, latent transforming
1748	19060	NM_021587	cc	1	growth factor beta binding protein 3
1749	17324	NM_021593	o,General		ESTs, Moderately similar to TYPE I
					IODOTHYRONINE DEIODINASE
				deiodinase, iodothyronine,	[R.norvegicus], deiodinase,
1750	19679	NM_021653	General	type I	iodothyronine, type I
					ESTs, Moderately similar to TYPE I IODOTHYRONINE DEIODINASE
				deiodinase, iodothyronine,	[R.norvegicus], deiodinase,
1750	19678	NM 021653	a,v,General	type I	iodothyronine, type I
			l		potassium channel, subfamily K,
					member 1 (TWIK-1), potassium
					channel, subfamily K, member 3
]					(TASK-1), potassium channel, subfamily K, member 6 (TWIK-2),
					potassium channel, subfamily K,
1751	19665	NM 021688	u,General		member 7

TABLE 8	HUMANI	HONOLOGUE AN	EKONTATOKI		Aity. Docket No. 44921-5039WO Doc. No. 1798897.1
Seq. ID No.	ldentifier	GeriBenik Ace./ Ref. Seq. 10 No.	Model Gode	Homologove Gene Name	Homologovs Elvstör Name
No.	Identifier	Ref. Seq. 1D. No.	Model Gode	Name	EST, Highly similar to GUANINE NUCLEOTIDE RELEASING PROTEIN [R.norvegicus], EST, Weakly similar to 2009427A guanine nucleotide-releasing protein [H.sapiens], ESTs, Highly similar to 2009427A guanine nucleotide-releasing protein [H.sapiens], ESTs, Weakly similar to S28407 guanine nucleotide-exchange activator CDC25 homolog - mouse [M.musculus], RAS protein-specific guanine nucleotide-releasing factor 1, RAS protein-specific guanine nucleotide-releasing factor 2, RIKEN cDNA 4921528G01 gene, Rap1 guanine-nucleotide-exchange factor directly activated by cAMP, Rattus norvegicus strain Wistar RAS guanine nucleotide-releasing factor 1 (Rasgrf1) mRNA, partial cds, guanine nucleotide-
1752	19667	NM_021690	 m		releasing factor 2 (specific for crk proto oncogene)
				prothymosin alpha, prothymosin, alpha (gene	ESTs, Highly similar to THYA_HUMAN PROTHYMOSIN ALPHA [H.sapiens], RIKEN cDNA 2610009E16 gene, prothymosin a14, prothymosin alpha, prothymosin, alpha (gene sequence
1754	22916	NM_021740	a	sequence 28)	28) CD14 antigen, ESTs, Weakly similar to CD14 RAT MONOCYTE
1755	19710	NM_021744	t		DIFFERENTIATION ANTIGEN CD14 PRECURSOR [R.norvegicus]
1755	19711	NM 021744	t		CD14 antigen, ESTs, Weakly similar to CD14 RAT MONOCYTE DIFFERENTIATION ANTIGEN CD14 PRECURSOR [R.norvegicus]
1756	19712	NM_021745	r		EST, Weakly similar to 138975 nuclear orphan receptor LXR-alpha [H.sapiens], ESTs, Moderately similar to JC4014 steroid hormone-nuclear receptor NER [H.sapiens], expressed sequence Al957360, nuclear receptor subfamily 1, group H, member 3, nuclear receptor subfamily 1, group H, member 4
1757	1962	NM_021750	j,k,y,z	1	Homo sapiens, clone MGC:18185
1757	19824	NM_021750	a,bb		Homo sapiens, clone MGC:18185 IMAGE:4155381, mRNA, complete cds, KIAA0251 hypothetical protein, RIKEN cDNA 1110027M19 gene, cysteine sulfinic acid decarboxylase- related protein 2, glutamic acid decarboxylase 2
1758	25198	NM_021754	h .		DKFZP566J153 protein, ESTs,
					Moderately similar to T17299 hypothetical protein DKFZp564H2171.1 [H.sapiens],
1758	20035	NM_021754	b,n,s,v,General	l	nucleolar protein NOP5/NOP58

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TABLE 3	E HUMANI	Hovorogne VX	SKÖTTATOKI		/Anny. Dooken No. 44921-5039W/O Doc. No. 17483977.1
Seq. ID No	ldenililer	Condonk Ace./ Ref. Seq. ID No.	Model Gode :	Homologovs Cana Name	Homologous Cluster Name
		200.257			EST, Weakly similar to LIS1 MOUSE PLATELET-ACTIVATING FACTOR ACETYLHYDROLASE IB ALPHA SUBUNIT [R.norvegicus], ESTs, Weakly similar to LIS1 MOUSE PLATELET-ACTIVATING FACTOR ACETYLHYDROLASE IB ALPHA SUBUNIT [R.norvegicus], Homo sapiens mRNA for FLJ00083 protein, partial cds, WD repeat domain 5, f-box and WD-40 domain protein 2, hypothetical protein, platelet-activating factor acetylhydrolase beta subunit (PAF-AH beta), platelet-activating factor acetylhydrolase, isoform 1b, beta1 subunit, platelet-activating factor acetylhydrolase, isoform Ib, alpha subunit (45kD), recombination protein
1759	17885	NM_021757	aa	(Arabidopsis)	REC14 ESTs, Weakly similar to COPP RAT COATOMER BETA' SUBUNIT [R.norvegicus], Mus musculus, Similar to RIKEN cDNA 1500041N16 gene, clone MGC:12066 IMAGE:3708188, mRNA, complete cds, RIKEN cDNA 2510040D07 gene, coatomer protein complex, subunit beta 2 (beta prime), expressed sequence AA408785, expressed sequence AI256832, expressed sequence C77982, glutamate rich WD repeat protein GRWD, platelet-activating factor acetylhydrolase, isoform 1b, beta1 subunit
1762	20161	NM 021836	cc,General		Jun-B oncogene, jun B proto- oncogene
					DKFZP586N1922 protein, ESTs, Moderately similar to S22695 restin [H.sapiens], ESTs, Weakly similar to T42734 cytoplasmic linker protein CLIP-115 - rat [R.norvegicus], RIKEN cDNA 1500005P14 gene, RIKEN cDNA 4631429H07 gene, RIKEN cDNA 5830409B12 gene, cytoplasmic linker 2, hypothetical protein FLJ20364, restin (Reed-Steinberg cell-espressed intermediate filament-associated protein), restin (Reed-Steinberg cell-expressed intermediate
1764	1203	NM_021997	k,z	cytoplasmic linker 2	filament-associated protein) EST, Moderately similar to PLM RAT PHOSPHOLEMMAN PRECURSOR [R.norvegicus], EST, Weakly similar to PLM_HUMAN PHOSPHOLEMMAN PRECURSOR [H.sapiens], FXYD domain-containing ion transport regulator 1, FXYD domain-containing ion transport regulator 1 (phospholemman), FXYD domain- containing ion transport regulator 3, FXYD domain-containing ion transport regulator 4, FXYD domain-containing ion transport regulator 6, FXYD
1765	23151	NM_022005	b	FXYD domain-containing ion transport regulator 6	domain-containing ion transport regulator 7, hypothetical protein MGC13186

PARIUS A		HOMOLOGUEAN	INATATIANS	7.3.02	
					Doc. No. 179889
Seg. [D. No.	ldentifier	Geneant Ace <i>l</i> Rof. Seq. ID No.	Model Code	Hamalogous Cana	Homologous Gluster Name
					EST, Moderately similar to HXK3_HUMAN HEXOKINASE TYF III [H.sapiens], EST, Weakly similar HXK2 MOUSE HEXOKINASE TYP [M.musculus], ESTs, Moderately similar to HXK2 MOUSE HEXOKINASE TYPE II [M.musculu ESTs, Weakly similar to HXK2 MOUSE HEXOKINASE TYPE II [M.musculus], hexokinase 2,
1767	17101	NM_022179	bb		hexokinase 3 (white cell) EST, Moderately similar to HXK3_HUMAN HEXOKINASE TYP III [H.sapiens], EST, Weakly simila HXK2 MOUSE HEXOKINASE TYP [M.musculus], ESTs, Moderately similar to HXK2 MOUSE HEXOKINASE TYPE II [M.musculus], ESTs, Weakly similar to HXK2 MOUSE HEXOKINASE TYPE II [M.musculus], hexokinase 2,
1767	17100	NM_022179	bb		hexokinase 3 (white cell) expressed sequence AA986699,
1768	20257	NM 022180	w,General		hepatic nuclear factor 4, hepatocyt nuclear factor 4, alpha
1768	25699	NM_022180	i		expressed sequence AA986699, hepatic nuclear factor 4, hepatocyt nuclear factor 4, alpha
1768	10860	NM_022180	k,x	topoisomerase (DNA) II alpha, topoisomerase (DNA) II alpha (170kD)	ESTs, Moderately similar to A4049 DNA topoisomerase [H.sapiens], ESTs, Moderately similar to TP2A MOUSE DNA TOPOISOMERASE ALPHA [M.musculus], topoisomera (DNA) II alpha, topoisomerase (DN beta, topoisomerase (DNA) II beta (180kD)
1770	20312	NM 022224	o		expressed sequence Al790318, expressed sequence Al836570, phosphotriesterase related
					WNT1 inducible signaling pathway protein 1, WNT1 inducible signaling pathway protein 2, WNT1 inducible signaling pathway protein 3,
1771	6585	NM_022266	d,p,cc		connective tissue growth factor ESTs, Highly similar to A23035 tub alpha chain [H.sapiens], tubulin alp 1, tubulin alpha 2, tubulin alpha 3, tubulin alpha 6, tubulin, alpha 2, tubulin, alpha 3, tubulin, alpha,
1772	17161	NM_022298	i,v,cc,General		ubiquitous ESTs, Highly similar to A23035 tub alpha chain [H.sapiens], tubulin alp 1, tubulin alpha 2, tubulin alpha 3, tubulin alpha 6, tubulin, alpha 2, tubulin, alpha 3, tubulin, alpha,
1772 1772	17162	NM_022298	u		ubiquitous ESTs, Highly similar to A23035 tub alpha chain [H.sapiens], tubulin alp 1, tubulin alpha 2, tubulin alpha 3, tubulin alpha 6, tubulin, alpha 2, tubulin, alpha 3, tubulin, alpha, ubiquitous

Mark State		HOMOLOGUE AN	NOTATIONS		Ally Docket No. 44924-5039W© Doc. No. 1798897.1
Seg. (D). No.	ldentifier	Cenbenk Acq./ Ref. Seq. ID No.	Model Code	Homologous Care Name	Homologous Cluster Neme
	Tables 4 to china manazioni are directifi	particle designation of the second of the se	Charles of the Charle		ESTs, Highly similar to A23035 tubulin alpha chain [H.sapiens], tubulin alpha 1, tubulin alpha 2, tubulin alpha 3, tubulin alpha 6, tubulin, alpha 2, tubulin, alpha 3, tubulin, alpha,
1772	17158	NM_022298	g		ubiquitous EST, Moderately similar to E Chain E,
1773	11454	NM_022381	i,aa,General		Human Pcn [H.sapiens], proliferating cell nuclear antigen
		NINA 000004	1.001		EST, Moderately similar to E Chain E, Human Pcn [H.sapiens], proliferating
1773	13480	NM_022381 NM_022390	I,General s	0.70.	cell nuclear antigen RIKEN cDNA 2610008L04 gene, quinoid dihydropteridine reductase
1774	15184	NM 022391	z	pituitary tumor- transforming 1	quinoia umyaraptename reductaso
1775	13104	NW_022591		nansoming i	EST, Highly similar to ISI1_HUMAN INSULIN-INDUCED PROTEIN 1 [H.sapiens], ESTs, Highly similar to INSULIN-INDUCED GROWTH RESPONSE PROTEIN CL-6 [R.norvegicus], Mus musculus, clone MGC:18904 IMAGE:4240711, mRNA, complete cds, insulin induced gene 1,
1776	22413	NM_022392	h		insulin induced protein 2 EST, Highly similar to ISI1_HUMAN
					INSULIN-INDUCED PROTEIN 1 [H.sapiens], ESTs, Highly similar to INSULIN-INDUCED GROWTH RESPONSE PROTEIN CL-6 [R.norvegicus], Mus musculus, clone MGC:18904 IMAGE:4240711, mRNA, complete cds, insulin induced gene 1,
1776	22414	NM_022392	n		insulin induced protein 2 C-type (calcium dependent,
1777	22499	NM 022393	t		carbohydrate recognition domain) lectin, superfamily member 10, C-type (calcium dependent, carbohydrate recognition domain) lectin, superfamily member 6, C-type (calcium dependent, carbohydrate-recognition domain) lectin, superfamily member 6, CD72 antigen, Mus musculus Ly-49Q mRNA for NK receptor Ly-49Q, complete cds, Mus musculus, Similar to macrophage galactose N-acetyl-galactosamine specific lectin, clone MGC:25983 IMAGE:4456238, mRNA, complete cds, RIKEN cDNA 4930572L20 gene, asialoglycoprotein receptor 1, macrophage lectin 2 (calcium dependent)
1777	22499	NM_022393	Į į		ESTs, Weakly similar to
1779	24537	им_022399	е		CALRETICULIN PRECURSOR [R.norvegicus], Homo sapiens, Similar to RIKEN cDNA 1700031L01 gene, clone MGC:26577 IMAGE:4822010, mRNA, complete cds, RIKEN cDNA 1700031L01 gene, RIKEN cDNA 6330586120 gene, calreticulin

TABLE	E HUMAN	HOWOROGAL WY	SMOTATIONS		Affly, Docket No. 44921-5039W0 Doc. No. 1793397.1
Scq. [D No.		Consent Ace.	Model Gode :*	Homologous Cene Name	Homologous Gluster Namo
and the second s	la land				ESTs, Weakly similar to CALRETICULIN PRECURSOR [R.norvegicus], Homo sapiens, Similar to RIKEN cDNA 1700031L01 gene, clone MGC:26577 IMAGE:4822010, mRNA, complete cds, RIKEN cDNA
1779	24539	NM_022399	у		1700031L01 gene, RIKEN cDNA 6330586l20 gene, calreticulin
					ESTs, Highly similar to 139161 dystonin isoform 2 [H.sapiens], ESTs, Weakly similar to T42725 actin binding protein ACF7, neural isoform 1 - mouse (fragment) [M.musculus], Leman coiled-coil protein, actin- crosslinking protein 7, expressed sequence AA591047, expressed sequence AW554249, plectin 1, intermediate filament binding protein, 500kD, serologically defined colon
1780	1141	NM_022401	o,General		cancer antigen 8 EST, Highly similar to R5HUP0 acidic
					ribosomal protein P0, cytosolic [H.sapiens], EST, Moderately similar to RLA0 RAT 60S ACIDIC RIBOSOMAL PROTEIN P0 [R.norvegicus], ESTs, Highly similar to R5HUP0 acidic ribosomal protein P0, cytosolic [H.sapiens], ESTs, Highly similar to RLA0 MOUSE 60S ACIDIC RIBOSOMAL PROTEIN P0 [M.musculus], RIKEN cDNA 2610012O22 gene, acidic ribosomal phosphoprotein P0, hypothetical protein IMAGE3455200, ribosomal
1781	1069	NM_022402	g		protein, large, P0 ESTs, Highly similar to FRHUL ferritin
1782	8211	NM_022500	j,n,s	ferritin light chain 1, ferritin, light polypeptide	light chain [H.sapiens], ESTs, Moderately similar to FRHUL ferritin light chain [H.sapiens], PRO0470 protein, RIKEN cDNA 4933416E14 gene, ferritin light chain 2, ferritin, light polypeptide ESTs, Highly similar to FRHUL ferritin
				ferritin light chain 1, ferritin,	light chain [H.sapiens], ESTs, Moderately similar to FRHUL ferritin light chain [H.sapiens], PRO0470 protein, RIKEN cDNA 4933416E14 gene, ferritin light chain 2, ferritin, light
1782	8212	NM_022500	n,s	light polypeptide cytochrome c oxidase subunit VIIa polypeptide 3 (liver), cytochrome c	polypeptide EST, Moderately similar to OSHU7L cytochrome-c oxidase [H.sapiens], cytochrome c oxidase subunit VIIa polypeptide 2 (liver), cytochrome c oxidase, subunit VIIa 1, cytochrome c
1783	6815	NM_022503	s	oxidase, subunit VIIa 3	oxidase, subunit VIIa 3 EST, Moderately similar to T08720 ribosomal protein L36 [H.sapiens], ESTs, Weakly similar to RL36_HUMAN 60S RIBOSOMAL PROTEIN L36 [H.sapiens], RIKEN
1784	4259	NM_022504	q,w	ribosomal protein L36	cDNA 1110038G14 gene, ribosomal protein L36

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	Ziáši .	HOMOLOGUE A)	ROTATIONS		Atty: Docket No. 44921-5088W Doc. No. 1793897
Seq. (D). Yo.	ldentifier	Cenbenk Ace <i>l</i> Rei. Seq. ID No.	Model Code	Homologous Care! Name	Homologous Cluster Name
1785	1611	NM 022509			ESTs, Weakly similar to SMN1_HUMAN SURVIVAL MOTOR NEURON PROTEIN 1 [H.sapiens], RIKEN cDNA 2410004J23 gene, expressed sequence Al849087, expressed sequence AW122398, splicing factor 30, survival of motor neuron-related, survival motor neuro survival motor neuron pseudogene, survival of motor neuron 1, telomeric survival of motor neuron 2, centromeric
					ESTs, Weakly similar to acyl-CoA dehydrogenase [R.norvegicus], RIKE cDNA 1300003O09 gene, RIKEN cDNA 2310016C19 gene, acetyl-Coenzyme A dehydrogenase, short chain, acyl-Coenzyme A dehydrogenase family, member 8, ac Coenzyme A dehydrogenase family, member 8, ac Toenzyme A dehydrogenase, C-2 to 3 short chain, hypothetical protein FLJ12592, hypothetical protein MGC5601, isovaleryl coenzyme A dehydrogenase
1786 1787	3026	NM_022512	y,z la		EST, Weakly similar to 60S RIBOSOMAL PROTEIN L27 [R.norvegicus], EST, Weakly similar S43505 ribosomal protein L27, cytosolic [H.sapiens], ribosomal protein L27
1787	3027	NM_022514	a,q,r,aa		EST, Weakly similar to 60S RIBOSOMAL PROTEIN L27 [R.norvegicus], EST, Weakly similar S43505 ribosomal protein L27, cytosolic [H.sapiens], ribosomal protein L27
1788	2696	NM_022515	a,d		ESTs, Highly similar to RL24_HUMA 60S RIBOSOMAL PROTEIN L24 [H.sapiens], RIKEN cDNA 0610008I gene, ribosomal protein L24
1788	2697	NM_022515	n,w,aa		ESTs, Highly similar to RL24_HUMA 60S RIBOSOMAL PROTEIN L24 [H.sapiens], RIKEN cDNA 0610008I gene, ribosomal protein L24
1789	3900	NM_022516	h	neural polypyrimidine tract binding protein, polypyrimidine tract binding protein	ESTs, Moderately similar to S15552 polypyrimidine tract-binding protein rat [R.norvegicus], ESTs, Weakly similar to S15552 polypyrimidine trabinding protein 1 - rat [R.norvegicus RIKEN cDNA 2810036L13 gene, expressed sequence AW107884, heterogeneous nuclear ribonucleoprotein L, neural polypyrimidine tract binding protein, polypyrimidine tract binding protein, polypyrimidine tract binding protein ADP-ribosylation factor 1, ADP-ribosylation factor 1, ADP-fibosylation factor domain protein 1, 64kD, ADP-ribosylation factor-like 1
1790	4151	NM 022518	o		ARF protein, Homo sapiens, Similar DKFZP727C091 protein, clone MGC:10677 IMAGE:3948445, mRN complete cds

TABLE	S HUMAN	HOMOLOGUE AN	NOTATIONS		Aijy, Dockei No. 44921-5039WO Doc. No. 1793397,1
Seg. [D No. :	Identifier	GenBank Acc./ Ref. Seq. ID No.	Model Gode, 🕭	Homologous Gana Namen	Homologous Cluster Name
Section 2 Control	Delivery and god to deliver with the	THE STATE OF THE S			ESTs, Highly similar to ORNITHINE AMINOTRANSFERASE PRECURSOR [R.norvegicus], RIKEN cDNA 1300019H02 gene, RIKEN cDNA 2900006B13 gene, hypothetical protein MGC15875, ornithine aminotransferase, ornithine
1791	4242	NM_022521	С		aminotransferase (gyrate atrophy) CD151 antigen, Homo sapiens cDNA
					FLJ14609 fis, clone NT2RP1000944, RIKEN cDNA 1110014F12 gene, RIKEN cDNA 1110031P12 gene, RIKEN cDNA 2700063A19 gene, transmembrane 4 superfamily member (tetraspan NET-2), transmembrane 4
1792	4412	NM_022523	0		superfamily member 6 ESTs, Weakly similar to A55046
1793	6641	NM_022533	General		plasmolipin - rat [R.norvegicus], ESTs, Weakly similar to PLLP_HUMAN PLASMOLIPIN [H.sapiens], Homo sapiens cDNA FLJ14787 fis, clone NT2RP4000878, highly similar to MYELOID UPREGULATED PROTEIN, Mus musculus, Similar to BENE protein, clone MGC:19097 IMAGE:4205488, mRNA, complete cds, RIKEN cDNA 2700018N07 gene, expressed sequence Al461653, myelin and lymphocyte protein; T-cell differentiation protein, plasmolipin
					ESTs, Moderately similar to A Chain A, Cyclophilin B Complexed With [H.sapiens], RIKEN cDNA 1110060010 gene, RIKEN cDNA 3732410E19 gene, RIKEN cDNA 4833408F11 gene, expressed sequence AA408962, expressed
					sequence AA553318, peptidylprolyl isomerase B, peptidylprolyl isomerase
1794	8097	NM_022536	а		B (cyclophilin B)
1795	8597	NM_022538	c,r,u		
1795	8598	NM_022538	u		EST, Highly similar to IM8B_MOUSE MITOCHONDRIAL IMPORT INNER MEMBRANE TRANSLOCASE SUBUNIT TIM8 B (DEAFNESS DYSTONIA PROTEIN 2 HOMOLOG) [R.norvegicus], translocase of inner mitochondrial membrane 8 (yeast) homolog B, translocase of inner mitochondrial membrane 8 homolog b
1796	9296	NM_022541	0	<u> </u>	(yeast) ESTs, Weakly similar to A Chain A,
	21063	NM_022585	h	ornithine decarboxylase antizyme inhibitor	Crystal Structure Ornithine Decarboxylase From Mouse, Truncated 37 Residues From The C- Terminus, To 1.6 Angstrom Resolution [M.musculus], ESTs, Weakly similar to ORNITHINE DECARBOXYLASE [R.norvegicus], Ornitine decarboxylase, ornithine decarboxylase 1, ornithine decarboxylase antizyme inhibitor, ornithine decarboxylase, structural, ornithine decarboxylase, structural,

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TABLE	S: HUMAN	HOMOLOGUE AL	SKIOTEATOKK		Aijy. Docket No. 44921-5139W0 Doc. No. 1799397.1
Seq. ID No.	(Identifier	Conbank Ace./ Ref. Seq. ID No.		Homologous Cene Name	Homologous Cluster Name
1799	20781	NM_022591	z		apoptotic protease activating factor, telomerase associated protein 1, telomerase-associated protein 1
1800	20803	NM_022592	n		RIKEN cDNA 4933401119 gene, hypothetical protein DKFZp434L1717, transketolase, transketolase (Wernicke Korsakoff syndrome), transketolase- like 1
1801	20925	NM_022594	q	enoyl Coenzyme A hydratase 1, peroxisomal, enoyl coenzyme A hydratase 1, peroxisomal	AU RNA-binding protein/enoyl- coenzyme A hydratase, EST, Weakly similar to ECH1_HUMAN DELTA3,5- DELTA2,4-DIENOYL-COA ISOMERASE PRECURSO [H.sapiens], RIKEN cDNA 2610009M20 gene, enoyl Coenzyme A hydratase 1, peroxisomal, enoyl coenzyme A hydratase 1, peroxisomal, expressed sequence AA617331
1802	20944	NM 022597	aa		cathepsin B, lipocalin 7, tubulointerstitial nephritis antigen
1602	20944	NW_022391			EST, Highly similar to OM25_RAT MITOCHONDRIAL OUTER MEMBRANE PROTEIN 25 (NPW16) [R.norvegicus], EST, Weakly similar to OM25_RAT MITOCHONDRIAL OUTER MEMBRANE PROTEIN 25 (NPW16) [R.norvegicus], Erbb2 interacting protein, discs, large homolog 4 (Drosophila), expressed sequence Al118201, hypothetical protein FLJ11271, synaptojanin 2
1803	21024	NM_022599	o,General		binding protein
1804	2250	NM_022643	General		H2B histone family, member D, H2B histone family, member G, H2B histone family, member K, RIKEN cDNA 2610022J01 gene, expressed sequence Al413321, expressed sequence R74621
1805	17567	NM_022672	a,y	ribosomal protein S14	EST, Moderately similar to 40S RIBOSOMAL PROTEIN S14 [R.norvegicus], EST, Weakly similar to A25220 ribosomal protein S14, cytosolic [H.sapiens], EST, Weakly similar to JE0129 ribosomal protein S14 - mouse [M.musculus], ESTs, Highly similar to A25220 ribosomal protein S14, cytosolic [H.sapiens], ribosomal protein S14
				H2A histone family,	EST, Weakly similar to histone H2A.F/Z variant [H.sapiens], ESTs, Highly similar to HISTONE H2A.Z [R.norvegicus], H2A histone family, member Z. RIKEN cDNA C530002L11
1806	17661	NM_022674	bb	member Z	gene, histone H2A.F/Z variant EST, Weakly similar to IPP1_HUMAN PROTEIN PHOSPHATASE INHIBITOR 1 [H.sapiens], ESTs, Weakly similar to IPP1_HUMAN PROTEIN PHOSPHATASE INHIBITOR 1 [H.sapiens], RIKEN cDNA 4930579P15 gene, protein phosphatase 1, regulatory (inhibitor) subunit 1A, protein phosphatase 1, regulatory (inhibitor) subunit 1B
1807	24563	NM_022676	b	regulatory (inhibitor) subunit 1A	(dopamine and cAMP regulated phosphoprotein, DARPP-32)

		HOMOLOGUEA			Ally, Docket No. 44921-503900 Doc. No. 1793397.
Seg. ID No.	් ල්ලාබ්බීල්	Cenenk Acel Reil Seg. ID No.	Model Code	Honologous Gene Name	Homologgus Cluster Name
1807	24564	NM_022676	b,x	protein phosphatase 1, regulatory (inhibitor) subunit 1A	EST, Weakly similar to IPP1_HUMAN PROTEIN PHOSPHATASE INHIBITOR 1 [H.sapiens], ESTs, Weakly similar to IPP1_HUMAN PROTEIN PHOSPHATASE INHIBITOR 1 [H.sapiens], RIKEN cDNA 4930579P15 gene, protein phosphatase 1, regulatory (inhibitor) subunit 1A, protein phosphatase 1, regulatory (inhibitor) subunit 1B (dopamine and cAMP regulated phosphoprotein, DARPP-32)
1808	20506	NM_022686	<u> </u>	<u> </u>	
1809	20508	NM_022688	9		EBNA-2 co-activator (100kD), ESTs, Moderately similar to l38968 100 kDa
1810	17586	NM_022694	k		coactivator [H.sapiens] DNA segment, Chr 7, Wayne State University 21, expressed, EST, Highly similar to S55915 ribosomal protein L28 [H.sapiens], ESTs, Highly similar to S55915 ribosomal protein L28
1811	17730	NM_022697	a		[H.sapiens], ribosomal protein L28 DNA segment, Chr 7, Wayne State
					University 21, expressed, EST, Highly similar to S55915 ribosomal protein L28 [H.sapiens], ESTs, Highly similar to S55915 ribosomal protein L28
1811	17729	NM_022697	q		[H.sapiens], ribosomal protein L28
1812	154	NM_022849	t	crp-ductin, deleted in malignant brain tumors 1	CD163 antigen, ESTs, Highly similar (138005 M130 antigen precursor, splict form 4 [H.sapiens], KIAA1822 protein apoptosis inhibitory 6, crp-ductin, deleted in malignant brain tumors 1, lectin, galactoside-binding, soluble, 3 binding protein, macrophage scavenger receptor 2, peptidylprolyl isomerase C-associated protein
1040	407	NM 002055			EST, Weakly similar to KC13 RAT CASEIN KINASE I, GAMMA 3 ISOFORM [R.norvegicus], ESTs, Highly similar to KC13 RAT CASEIN KINASE I, GAMMA 3 ISOFORM [R.norvegicus], ESTs, Weakly similar to casein kinase [M.musculus], RIKEN cDNA 2610208K14 gene, RIKEN cDNA 3300002K07 gene, casein kinase 1, alpha 1, casein kinase 1, delta, casein kinase 1, epsilon, caseir kinase 1, gamma 1, casein kinase 1,
1813	127	NM_022855	h		gamma 3 ESTs, Highly similar to FXD3_HUMAI FORKHEAD BOX PROTEIN D3
					[H.sapiens], ESTs, Weakly similar to FXD3_HUMAN FORKHEAD BOX PROTEIN D3 [H.sapiens], ESTs, Weakly similar to HFH1 RAT HEPATOCYTE NUCLEAR FACTOR FORKHEAD HOMOLOG 1 [R.norvegicus], HNF-3/forkhead homolog 1 like, expressed sequence
					Al385632, forkhead box D1, forkhead box D2, forkhead box D3, winged
1814 1816	152 18101	NM_022858 NM_022948	j z		Al385632, forkhead box D1, forkhead

TABUE®	E'HWMAN	HOMOLOGUE AK	NOTATIONS		Ally, Docket No. 44921-5039WO Doc. No. 1793397.1
Seq. (10) (No.	Identifier	GenBank Acc./ Ref. Seq. ID No.	Model Gode	Homologous Gene Name	Homologous Cluster Name
hand of consenses.					CAT56 protein, EST, Highly similar to CA13_HUMAN COLLAGEN ALPHA 1(III) CHAIN PRECURSOR [H.sapiens], EST, Weakly similar to A28996 proline-rich protein M14 precursor - mouse [M.musculus], EST, Weakly similar to JE0291 FB19 protein [H.sapiens], collagen, type III, alpha 1 (Ehlers-Danlos syndrome type IV, autosomal dominant), procollagen, type III, alpha 1, proline rich protein,
1817	21491	NM 022951	w		proline rich protein 2, protein phosphatase 1, regulatory subunit 10
					ESTs, Moderately similar to S57219 phosphatidylinositol 3-kinase [H.sapiens], FK506 binding protein 12-rapamycin associated protein 1, Homo sapiens cDNA FLJ12591 fis, clone NT2RM4001313, moderately similar to PHOSPHATIDYLINOSITOL 3-KINASE VPS34-LIKE (EC 2.7.1.137), Homo sapiens cDNA FLJ14331 fis, clone PLACE4000320, RIKEN cDNA 2410099E07 gene, phosphatidylinositol 3-kinase catalytic delta polypeptide, phosphatidylinositol 3-kinase, catalytic, beta polypeptide, phosphoinositide-3-kinase, class 3, rapamycin and FKBP12 target-1
1818	15742	NM_022958	у		protein
1819	9286	NM_023027	t,w		ELAV (embryonic lethal, abnormal vision, Drosophila)-like 4 (Hu antigen D), ESTs, Highly similar to HUD RAT PARANEOPLASTIC ENCEPHALOMYELITIS ANTIGEN HUD HOMOLOG [R.norvegicus], ESTs, Moderately similar to HUD RAT PARANEOPLASTIC ENCEPHALOMYELITIS ANTIGEN HUD HOMOLOG [R.norvegicus], ESTs, Moderately similar to POLYADENYLATE-BINDING PROTEIN 1 [M.musculus], Mus musculus adult male testis cDNA, RIKEN full-length enriched library, clone:4933407N23, full insert sequence, RIKEN cDNA 4932702K14 gene, poly A binding protein, cytoplasmic 1
1819	9286	NM_023027	It,w		RIKEN cDNA 2610208K14 gene,
1820	23215	NM 023102	z		VRK3 for vaccinia related kinase 3, casein kinase 1, alpha 1, casein kinase 1, delta, casein kinase 1, epsilon, casein kinase 1, gamma 2
				CCAAT/enhancer binding	CCAAT/enhancer binding protein
1821	21238	NM_024125	cc,General	protein (C/EBP), beta CCAAT/enhancer binding protein (C/EBP), beta	(C/EBP), beta CCAAT/enhancer binding protein (C/EBP), beta
1821	21239	NM_024125	cc,General	growth arrest and DNA- damage-inducible 45 alpha, growth arrest and DNA-damage-inducible,	growth arrest and DNA-damage- inducible 45 alpha, growth arrest and DNA-damage-inducible 45 beta, growth arrest and DNA-damage-
1822	353	NM_024127	i,n,General	alpha	inducible, alpha

TABLE	K IHUMAN	HOMOLOGUE AK	RIOTATIONS		:: Ally. Docket No. 44924-5039WO Doc. No. 1793397.1
Seq.(D) No.	refillmebl	ConBank Ace. Ref. Seq. 10 No.	Model Gode	Homologous Cone Name	Homologous Gluster Name
1822	354	NM 024127	i,n,General	growth arrest and DNA- damage-inducible 45 alpha, growth arrest and DNA-damage-inducible, alpha	growth arrest and DNA-damage- inducible 45 alpha, growth arrest and DNA-damage-inducible 45 beta, growth arrest and DNA-damage- inducible, alpha
1822	352	NM 024127	h,General	growth arrest and DNA- damage-inducible 45 alpha, growth arrest and DNA-damage-inducible, alpha	growth arrest and DNA-damage- inducible 45 alpha, growth arrest and DNA-damage-inducible 45 beta, growth arrest and DNA-damage- inducible, alpha
1823	17227	NM 024131	x	D-dopachrome tautomerase	D-dopachrome tautomerase, expressed sequence C78655
1824	1598	NM 024134	1	DNA-damage inducible transcript 3, DNA-damage-inducible transcript 3	DNA-damage inducible transcript 3, EST, Moderately similar to GA15_HUMAN GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE PROTEIN GADD153 [H.sapiens], myozenin
1825	1162	NM_024153	d	inducible transcript o	ferredoxin reductase
1826	7863	NM_024156	С		ATPase, H+ transporting, lysosomal (vacuolar proton pump) 16kD, ATPase, H+ transporting, lysosomal (vacuolar proton pump) 21kD, ESTs, Weakly similar to VACUOLAR ATP SYNTHASE 16 KD PROTEOLIPID SUBUNIT [R.norvegicus], Mus musculus, Similar to ATPase, H+ transporting, lysosomal (vacuolar proton pump) 21kD, clone MGC:6568 IMAGE:2812497, mRNA, complete cds EST, Weakly similar to A29154 complement factor I [H.sapiens], I factor (complement), RIKEN cDNA 1300008A22 gene, complement component factor i, protease, serine, 7
1827	22079	NM 024157	×		(enterokinase), suppression of tumorigenicity 14 (colon carcinoma), transmembrane protease, serine 2
1828	16476	NM_024162	General	fatty acid binding protein 3, muscle and heart, fatty acid binding protein 3, muscle and heart (mammary-derived growth inhibitor)	EST, Moderately similar to FABH MOUSE FATTY ACID-BINDING PROTEIN, HEART [M.musculus], ESTs, Highly similar to PC4011 fatty acid-binding protein - mouse [M.musculus], fatty acid binding protein 3, muscle and heart, fatty acid binding protein 3, muscle and heart (mammary-derived growth inhibitor), fatty acid binding protein 3, pseudogene 2 EST, Moderately similar to A27077 dnaK-type molecular chaperone
1829	17765	NM_024351	b.s.v	heat shock 70kD protein 8	[H.sapiens], EST, Weakly similar to A27077 dnaK-type molecular chaperone [H.sapiens], EST, Weakly similar to A45935 dnaK-type molecular chaperone hsc70 - mouse [M.musculus], heat shock 70kD protein 8, hypothetical protein MGC4859 similar to HSPA8

TABLE 3	HUMAN	HOWOTOGAEVK	NOTATIONS		Aity. Docket No. 44921-5039000 Doc. No. 1793897.1
Sco. ID දූරු	ldentifier	Conbook Ace./ Ref. Seq. ID No.	Model Gode	Homologous Gene Name	Homologous Cluster Name
					bHLH factor Hes4, basic helix-loophelix domain containing, class B, 2, basic helix-loop-helix domain containing, class B, 3, hairy (Drosophila)-homolog, hairy and enhancer of split 1, (Drosophila), hairy and enhancer of split 6, (Drosophila), hairy/enhancer-of-split related with YRPW motif 1, hairy/enhancer-of-split related with YRPW motif 2, likely ortholog of mouse Hes6 neuronal
1830	8879	NM_024360	h		differentiation gene ANM1_HUMAN PROTEIN ARGININE
1831	20772	NM_024363	×	HMT1 (hnRNP methyltransferase, S. cerevisiae)-like 2, heterogeneous nuclear ribonucleoproteins methyltransferase-like 2 (S. cerevisiae)	N-METHYLTRANSFERASE 1 [H.sapiens], EST, Weakly similar to ANM1_HUMAN PROTEIN ARGININE N-METHYLTRANSFERASE 1 [H.sapiens], ESTS, Moderately similar to ANM1_MOUSE PROTEIN ARGININE N- METHYLTRANSFERASE 1 [M.musculus], HMT1 (hnRNP methyltransferase, S. cerevisiae)-like 2, HMT1 (hnRNP methyltransferase, S. cerevisiae)-like 3, Homo sapiens cDNA: FLJ23133 fis, clone LNG08560, RIKEN cDNA 2410018A17 gene, coactivator-associated arginine methyltransferase-1, expressed sequence AW214366, heterogeneous nuclear ribonucleoproteins methyltransferase-like 2 (S. cerevisiae), heterogeneous nuclear ribonucleoproteins methyltransferase- like 2 (S. cerevisiae), related sequence, hypothetical protein FLJ10559
1001	20.72				3-hydroxy-3-methylglutaryl-Coenzyme A lyase, 3-hydroxymethyl-3-
1832	2812	NM_024386	С		methylglutaryl-Coenzyme A Iyase (hydroxymethylglutaricaciduria), Homo sapiens clone 24959 mRNA sequence partial cds, hypothetical protein
1833	335	NM_024387	j,y		RIKEN cDNA 2700048O17 gene, heme oxygenase (decycling) 2
1834	21	NM 024388	cc		nuclear receptor subfamily 4, group A, member 1
1834	22	NM 024388	СС		nuclear receptor subfamily 4, group A, member 1

TABLE 3	HUMAN	HOMOLOGUE AN	NOTATIONS -		Ally. Docket No. 44921-5039W0
Social de	H.40-36	BoxBox Acc # 1		M	Dos. No. 1793397.1 I
Sed: In. No:	ldentifier.	ConBank Ace. Ref. Seq. ID No.	Model Gode	Name Name	Homologous Avster Name
	See a see a see a see a see a see a see a see a see a see a see a see a see a see a see a see a see a see a se	General Annual Control of the Contro			Homo sapiens cDNA FLJ13261 fis, clone OVARC1000885, weakly similar to OXIDOREDUCTASE UCPA (EC 1), Human DNA sequence from clone 1068E13 on chromosome 20p11.21-12.3. Contains two putative novel genes, the gene for a novel protein similar to bovine SCP2 (Sterol Carrier Protein 2) and part of HSD17B4 (hydroxysteroid (17-beta) dehydrogenase 4), an EEF1A1 (eukaryotic translation elongation factor 1 alpha 1) pseudogene, ESTs, STSs and GSSs, RIKEN cDNA 1110029G07 gene, RIKEN cDNA 1700010M22 gene, RIKEN cDNA 1810026B04 gene, RIKEN cDNA 2610207116 gene, expressed sequence AW208803, hydroxysteroid (17-beta) dehydrogenase 4, hydroxysteroid 17-beta dehydrogenase 4, hypothetical protein
					MGC10940, oxidoreductase UCPA, retinal short-chain
1836	9929	NM_024392	f		dehydrogenase/reductase 1 ATP-binding cassette, sub-family A
4007	2502	NIM 024206			(ABC1), member 2, ATP-binding cassette, sub-family A (ABC1), member 3, ATP-binding cassette, sub-family A (ABC1), member 4, ATP-binding cassette, sub-family A (ABC1), member 7, ATP-binding cassette, sub-family A (ABC1), member 8, ESTs, Weakly similar to ABC2 MOUSE ATP-BINDING CASSETTE TRANSPORTER 2 [M.musculus], Homo sapiens mRNA for KIAA1888 protein, partial cds, hypothetical protein FLJ14297, hypothetical protein
1837	3582	NM_024396	aa		PRO2543 DNA segment, Chr 9, ERATO Doi 85,
1020	10003	NIM 024209	0.05.03		expressed, Homo sapiens aconitase precursor (ACON) mRNA, nuclear gene encoding mitochondrial protein, partial cds, RIKEN cDNA 5031409G22 gene, aconitase 1, aconitase 1, soluble, aconitase 2, mitochondrial, iron-responsive element-binding
1838	19993	NM_024398	e,p,s,aa		protein Homo sapiens, RIKEN cDNA
1839	10789	NM_024399	0		0610006H10 gene, clone MGC:17267 IMAGE:4155233, mRNA, complete cds, Homo sapiens, Similar to RIKEN cDNA 0610006H10 gene, clone MGC:9740 IMAGE:3853707, mRNA, complete cds, aspartoacylase (aminoacylase 2, Canavan disease)

TABLE 8	HUMAN)	HOMOLOGUE AV	- SKOITATOKÚ		Aily, Dockei No. 44921-5039W0 Doc. No. 1793397.1
Seq. ID No.	ldentifier	Condent Acel Ref. Seq. ID No.	Model Gode	Hemologove Cene Name	Homologous Guster Name
Section 1921 March 1970					ESTs, Weakly similar to T47158 hypothetical protein DKFZp762C1110.1 [H.sapiens], Mus musculus papilin mRNA, complete cds
					Mus musculus, Similar to a disintegrin and metalloproteinase with thrombospondin motifs 1 (ADAMTS-1).
					cione IMAGE:3491991, mRNA, partial cds, RIKEN cDNA 6720426B09 gene, RIKEN cDNA A930008K15 gene, a
					disintegrin-like and metalloprotease (reprolysin type) with thrombospondin type 1 motif, 1, a disintegrin-like and
					metalloprotease (reprolysin type) with thrombospondin type 1 motif, 4, a disintegrin-like and metalloprotease
					(reprolysin type) with thrombospondin type 1 motif, 8, a disintegrin-like and metalloprotease (reprolysin type) with
1840	22626	NM_024400	cc,General		thrombospondin type 1 motif, 9 EST, Weakly similar to ATF4_HUMAN CYCLIC-AMP-DEPENDENT TRANSCRIPTION FACTOR ATE 4
					TRANSCRIPTION FACTOR ATF-4 [H.sapiens], activating transcription factor 4 (tax-responsive enhancer element B67), activating transcription
1841	13633	NM_024403	g,General		factor 5 EST, Weakly similar to ATF4_HUMAN
					CYCLIC-AMP-DEPENDENT TRANSCRIPTION FACTOR ATF-4 [H.sapiens], activating transcription factor 4 (tax-responsive enhancer
1841	13634	NM_024403	g,General		element B67), activating transcription factor 5
					ESTs, Moderately similar to HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEIN C
					[R.norvegicus], ESTs, Weakly similar to A44192 heterogeneous nuclear ribonucleoprotein C-like protein
					[H.sapiens], Mus musculus high- glycine/tyrosine protein type I E5 mRNA, complete cds, expressed sequence C85084, heterogeneous
:					nuclear ribonucleoprotein A/B, heterogeneous nuclear ribonucleoprotein D, heterogeneous
1842	23387	NM_024404	b,General		nuclear ribonucleoprotein D (AU-rich element RNA-binding protein 1, 37kD) EST, Weakly similar to SYHUAL 5-
					aminolevulinate synthase [H.sapiens], ESTs, Moderately similar to 5- AMINOLEVULINIC ACID SYNTHASE
					MITOCHONDRIAL PRECURSOR, NONSPECIFIC [R.norvegicus], aminolevulinate, delta-, synthase 1,
					aminolevulinic acid synthase 1, aminolevulinic acid synthase 2, erythroid, serine palmitoyltransferase,
1843	21038	NM 024484	h	aminolevulinate, delta-, synthase 1, aminolevulinic acid synthase 1	long chain base subunit 1, serine palmitoyltransferase, long chain base subunit 2

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Seq. [D] No.] []	ldentifier	GenBank Acc./ Ref. Seq. ID No.	(1000) (1000)	Homologous Cene Name	Homologous Gluster Name
					GSHG_MOUSE GLUTATHIONE PEROXIDASE-GASTROINTESTINAL (GSHPX-GI) [M.musculus], ESTs, Weakly similar to GSHC RAT GLUTATHIONE PEROXIDASE [R.norvegicus], glutathione peroxidase
1844	1853	NM_030826	s	glutathione peroxidase 1	1, glutathione peroxidase 2 (gastrointestinal)
				low density lipoprotein receptor-related protein 2, low density lipoprotein-	EST, Highly similar to LRP2 RAT LOW-DENSITY LIPOPROTEIN RECEPTOR-RELATED PROTEIN 2 PRECURSOR [R.norvegicus], EST, Moderately similar to S25111 alpha-2-macroglobulin receptor precursor -mouse [M.musculus], ESTs, Highly similar to S02392 alpha-2-macroglobulin receptor precursor [H.sapiens], ESTs, Highly similar to S25111 alpha-2-macroglobulin receptor precursor eceptor precursor - mouse [M.musculus], ESTs, Weakly similar to I53413 calcium sensor protein [H.sapiens], expressed sequence C79313, low density lipoprotein receptor-related protein 2, low density lipoprotein 1, low density lipoprotein receptor-related protein 1, low density lipoprotein 1, low density lipoprotein 1, low density lipoprotein-related protein 1, low density lipoprotein-related
	15111	NM_030827	e,General	low density lipoprotein receptor-related protein 2, low density lipoprotein-related protein 2	lipoprotein-related protein 2 EST, Highly similar to LRP2 RAT LOW- DENSITY LIPOPROTEIN RECEPTOR- RELATED PROTEIN 2 PRECURSOR [R.norvegicus], EST, Moderately similar to S25111 alpha-2- macroglobulin receptor precursor - mouse [M.musculus], ESTs, Highly similar to S02392 alpha-2- macroglobulin receptor precursor [H.sapiens], ESTs, Highly similar to S25111 alpha-2-macroglobulin receptor precursor - mouse [M.musculus], ESTs, Weakly similar to I53413 calcium sensor protein [H.sapiens], expressed sequence C79313, low density lipoprotein receptor-related protein 1, low density lipoprotein receptor-related protein 2, low density lipoprotein 18 (deleted in tumors), low density lipoprotein-related protein 2

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Seq. (D No.	lidendifier	Consent Ace./ Rof. Seq. ID No.	Model Gode	Homologous Cane Name	Homologous Gluster Namo
1845	15110	NM_030827	General	low density lipoprotein receptor-related protein 2, low density lipoprotein- related protein 2	EST, Highly similar to LRP2 RAT LOW-DENSITY LIPOPROTEIN RECEPTOR-RELATED PROTEIN 2 PRECURSOR [R.norvegicus], EST, Moderately similar to \$25111 alpha-2-macroglobulin receptor precursor -mouse [M.musculus], ESTs, Highly similar to \$02392 alpha-2-macroglobulin receptor precursor [H.sapiens], ESTs, Highly similar to \$25111 alpha-2-macroglobulin receptor precursor - mouse [M.musculus], ESTs, Weakly similar to 153413 calcium sensor protein [H.sapiens], expressed sequence C79313, low density lipoprotein receptor-related protein 1, low density lipoprotein receptor-related protein 2, low density lipoprotein-related protein 1B (deleted in tumors), low density lipoprotein-related protein 2 solute carrier family 21 (organic anion
					solute carrier family 21 (organic anion transporter), member 1, solute carrier family 21 (organic anion transporter), member 10, solute carrier family 21 (organic anion transporter), member 14, solute carrier family 21 (organic anion transporter), member 3, solute carrier family 21 (organic anion transporter)
1846	808	NM_030837	k,m		transporter), member 6 ESTs, Weakly similar to I65309
1847	4057	NM 030844	k		autoantigen p69 - rat [R.norvegicus], Homo sapiens ALS2CR15 mRNA, partial cds, RIKEN cDNA 1700030B17 gene, islet cell autoantigen 1 (69kD), islet cell autoantigen 1, 69 kDa
1047	4037	NW_030044			GRO1 oncogene, GRO2 oncogene, Rattus norvegicus CXC chemokine RTCK1 (Rtck1) mRNA, complete cds, interleukin 8, platelet factor 4, pro- platelet basic protein, pro-platelet basic protein (includes platelet basic protein, beta-thromboglobulin, connective tissue-activating peptide III,
1848	1221	NM_030845	t		neutrophil-activating peptide-2) ESTs, Weakly similar to PERIPHERAL MYELIN PROTEIN 22 [R.norvegicus], Peripheral myelin protein, epithelial membrane protein 3, peripheral myelin protein 22, peripheral myelin protein,
1849	21509	NM_030847	x		22 kDa EST, Highly similar to PDK2 RAT [R.norvegicus], expressed sequence
				pyruvate dehydrogenase	Al035637, pyruvate dehydrogenase 2, pyruvate dehydrogenase kinase,
1850	1928	NM_030872	v	kinase, isoenzyme 2	isoenzyme 2 EST, Weakly similar to A Chain A, Human Platelet Profilin Complexed With The L-Pro10 Peptide (SUB 3-140 [H.sapiens], ESTs, Weakly similar to profilin [R.norvegicus], RIKEN cDNA
1851	17342	NM_030873	u		1700012P12 gene, profilin, profilin 1, profilin 2

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TABLE	BE HUMAN	HOMOLOGUEAN	NOTATIONS		Aity, Doctot No. 44921-5039700 Doc. No. 1793397.1
Seq. ID No.	ldentifler	ConBank Ace./ Ref. Seq. ID No.	Model Code	Memologous Gene Memo	Homologous Cluster Name
	11 (2111/2/18/00)	V			G protein-coupled receptor 15, G- protein coupled receptor SALPR; somatostatin and angiotensin-like peptide receptor, angiotensin receptor
1852	24648	NM 030985	u		1, expressed sequence AI551199
1852	25453	NM_030985	General		
					ESTs, Weakly similar to GBB1 RAT GUANINE NUCLEOTIDE-BINDING PROTEIN G(I)/G(S)/G(T) BETA SUBUNIT 1 [R.norvegicus], Mus musculus, clone MGC:7934 IMAGE:3583848, mRNA, complete cds, RIKEN cDNA 5930415H02 gene, constitutive photomorphogenic protein 1 (Arabidopsis), guanine nucleotide binding protein (G protein), beta polypeptide 1, guanine nucleotide binding protein beta subunit 4, guanine nucleotide binding protein beta subunit 4, guanine nucleotide binding protein, beta 1, guanine nucleotide binding protein, beta 4, similar to constitutive
1853	21802	NM 030987	h		photomorphogenic protein 1 (Arabidopsis)
					EST, Weakly similar to ALCOHOL DEHYDROGENASE [R.norvegicus], RIKEN cDNA 2310005E10 gene, aldoketo reductase family 1, member A1 (aldehyde reductase), aldo-keto reductase family 1, member A4 (aldehyde reductase), aldo-keto
1854	23109	NM_031000	f,s,z		reductase family 1, member E1 4-aminobutyrate aminotransferase,
1855	134	NM 031003	la,u		RIKEN cDNA 1300019H02 gene, RIKEN cDNA 2900006B13 gene, ornithine aminotransferase
1856	25461	NM 031009	0		
1857	1845	NM 031010	t		ARACHIDONATE 12- LIPOXYGENASE [R.norvegicus], arachidonate 12-lipoxygenase, arachidonate 12-lipoxygenase pseudogene 2, arachidonate 15- lipoxygenase, expressed sequence AW259591
1857	25517	NM 031010	c,t		ARACHIDONATE 12- LIPOXYGENASE [R.norvegicus], arachidonate 12-lipoxygenase, arachidonate 12-lipoxygenase pseudogene 2, arachidonate 15- lipoxygenase, expressed sequence AW259591
1858	16562	NM_031020	f		ESTs, Weakly similar to MK14 RAT MITOGEN-ACTIVATED PROTEIN KINASE 14 [R.norvegicus], mitogen activated protein kinase 14, mitogen-activated protein kinase 11, mitogen-activated protein kinase 14
1000	10302	14141_001020	<u>'</u>	 	casein kinase 2, beta polypeptide,
1859	1480	NM_031021	f		casein kinase II, beta subunit

TABLE	Se ituman	HOMOTOGALE VA	EMOTRATON		Aliy, Docket No. 44921-5039WO Doc. No. 1793897.1
Seq. ID No.	ldentifier	Conbank Ace./ Ref. Seg. ID No.	Model Gode	Homologous Cene Namo	Homologous Auster Name
1860	1719	NM_031024	n	drebrin 1	ESTs, Moderately similar to T30989 serine/threonine protein kinase NIK - mouse [M.musculus], RIKEN cDNA 1500031A17 gene, drebrin 1, drebrinlike, hypothetical protein FLJ13154, mitogen-activated protein kinase kinase kinase kinase kinase kinase kinase kinase kinase kinase kinase kinase kinase kinase containing protein HIP-55
					DnaJ (Hsp40) homolog, subfamily B, member 6, EST, Moderately similar to NEK1 MOUSE SERINE/THREONINE-PROTEIN KINASE NEK1 [M.musculus], EST, Moderately similar to T31096 cyclin G-associated kinase GAK - rat [R.norvegicus], EST, Weakly similar to T31096 cyclin G-associated kinase GAK - rat [R.norvegicus], EST, Moderately similar to NEK1 MOUSE SERINE/THREONINE-PROTEIN KINASE NEK1 [M.musculus], KIAA1048 protein, Mus musculus, Similar to cyclin G associated kinase, clone IMAGE:3487931, mRNA, partial cds, NIMA (never in mitosis gene a)-related expressed kinase 1, RIKEN cDNA 4632401F23 gene, cyclin G associated kinase, hypothetical protein DKFZp434P0116, serine/threonine
1861	1350	NM_031030	h	1.	kinase 16 ESTs, Highly similar to GLYCINE
1862	16775	NM 031031	General		AMIDINOTRANSFERASE PRECURSOR [R.norvegicus], RIKEN cDNA 1810003P21 gene, glycine amidinotransferase (L-arginine:glycine amidinotransferase)
1863	691	NM_031034	w		ESTs, Highly similar to GB12 RAT GUANINE NUCLEOTIDE-BINDING PROTEIN, ALPHA-12 SUBUNIT [R.norvegicus], guanine nucleotide binding protein (G protein) alpha 12, guanine nucleotide binding protein (G protein), alpha 13, guanine nucleotide binding protein, alpha 12, guanine nucleotide binding protein, alpha 12, guanine nucleotide binding protein, alpha 13
1864	15886	NM_031035	z		guanine nucleotide binding protein (G protein), alpha inhibiting activity polypeptide 2, guanine nucleotide binding protein, alpha inhibiting 2, guanine nucleotide binding protein, alpha inhibiting 3
1866	3608	NM 031044	k,General		Homo sapiens, Similar to histamine N- methyltransferase, clone MGC:14500 IMAGE:4249496, mRNA, complete cds, expressed sequence Al788969, histamine N-methyltransferase
			,		Homo sapiens, Similar to histamine N- methyltransferase, clone MGC:14500 IMAGE:4249496, mRNA, complete
1866	3610	NM_031044	d,General		cds, expressed sequence Al788969, histamine N-methyltransferase

TABLES	HUMAN	HOMOLOGUE AN	ROTATIONS		Atty, Doctot No. 44921-5039WO Doc, No. 1793897.1
Seg. (D No.	ndendifler	Consent Acel Ref. Seg. ID No.	Model Gode	Homologous Cene Name	Homologous Gluster Name
					EST, Highly similar to C Chain C, Macrophage Migration Inhibitory Factor [H.sapiens], EST, Moderately similar to C Chain C, Macrophage Migration Inhibitory Factor [H.sapiens], EST, Moderately similar to MIF RAT MACROPHAGE MIGRATION INHIBITORY FACTOR [R.norvegicus], macrophage migration inhibitory factor, macrophage migration inhibitory factor
1867	15137	NM_031051	s		(glycosylation-inhibiting factor) matrix metalloproteinase 14
1868	514	NM_031056	General		(membrane-inserted), matrix metalloproteinase 19, vitronectin
					RIKEN cDNA 1110038l05 gene, aldehyde dehydrogenase family 1, subfamily A4, expressed sequence Al427784, hypothetical protein FLJ23189, methylmalonate-
1869	17269	NM_031057	General		semialdehyde dehydrogenase EST, Moderately similar to R10A
4070	44040	NW 024055			MOUSE 60S RIBOSOMAL PROTEIN L10A [M.musculus], EST, Weakly similar to 60S RIBOSOMAL PROTEIN L10A [R.norvegicus], ESTs, Highly similar to R10A_HUMAN 60S RIBOSOMAL PROTEIN L10A [H.sapiens], ribosomal protein L10A, ribosomal protein L10a
1870	11849	NM_031065	a.	nucleoporin 98,	EST, Highly similar to NU98 RAT NUCLEAR PORE COMPLEX PROTEIN NUP98 [R.norvegicus], ESTs, Weakly similar to NU98 RAT NUCLEAR PORE COMPLEX PROTEIN NUP98 [R.norvegicus], RIKEN cDNA 0610038H21 gene, RIKEN cDNA 4930432K09 gene, RIKEN cDNA 5430432N15 gene, melanoma antigen, family D, 3, nucleoporin 98kD, plasma membrane
1871	1855	NM_031074	h	nucleoporin 98kD	associated protein, S3-12, trophinin
1872	4683	NM 031083	d		phosphatidylinositol 3-kinase, catalytic, alpha polypeptide, phosphatidylinositol 4-kinase, catalytic, beta polypeptide, phosphoinositide-3-kinase, catalytic, gamma polypeptide
					EST, Weakly similar to RALA MOUSE RAS-RELATED PROTEIN RAL-A [M.musculus], Homo sapiens cDNA: FLJ23197 fis, clone REC00917, RIKEN cDNA 1110065D03 gene, v-ral simian leukemia viral oncogene homolog A (ras related), v-ral simian leukemia viral oncogene homolog B
1873	15202	NM_031093	a a,n		(ras related) EST, Weakly similar to RALA MOUSE RAS-RELATED PROTEIN RAL-A [M.musculus], Homo sapiens cDNA: FLJ23197 fis, clone REC00917, RIKEN cDNA 1110065D03 gene, v-ral simian leukemia viral oncogene homolog A (ras related), v-ral simian leukemia viral oncogene homolog B (ras related)

TABLE &	KUMANI	HOMOLOGAE VIX	NOTATIONS		Aijy, Docksi No. 44921-5039WO Doc. No. 1798897.1
Seg. ID No.	dentiller	ConBank Ace./ Ref. Seq./D No.	Model Code	Homologous Cane Name	Homologovis Cluster Name
		NIM 024000			ESTs, Weakly similar to S55912 ribosomal protein L5, cytosolic
1874	12639	NM_031099	aa		[H.sapiens], ribosomal protein L5 EST, Moderately similar to 60S RIBOSOMAL PROTEIN L10 [M.musculus], EST, Moderately similar to 60S RIBOSOMAL PROTEIN L10 [R.norvegicus], ESTs, Highly similar to A42735 ribosomal protein L10, cytosolic [H.sapiens], Homo sapiens, Similar to ribosomal protein L10, clone MGC:22634 IMAGE:3935452, mRNA, complete cds, Human DNA sequence from clone RP3-334F4 on chromosome 6 Contains ESTs, STSs and GSSs. Contains a LAMR1 (laminin receptor 1, ribosomal protein SA) pseudogene and an RPL10 (ribosomal protein L10) pseudogene, Mouse 24.6 kda protein mRNA, complete cds,
1875	20812	NM_031100	a		ribosomal protein L10 ESTs, Weakly similar to 60S
1876	16938	NM 031103	w		RIBOSOMAL PROTEIN L19 [R.norvegicus], Homo sapiens mRNA; cDNA DKFZp434D115 (from clone DKFZp434D115), ribosomal protein L19
					ESTs, Highly similar to JC2120 heparin-binding protein 15 [H.sapiens], ESTs, Moderately similar to RL22 RAT 60S RIBOSOMAL PROTEIN L22 [R.norvegicus], Homo sapiens mRNA; cDNA DKFZp586E0524 (from clone DKFZp586E0524), Human DNA sequence from clone 581F12 on chromosome Xq21. Contains Eukaryotic Translation Initiation Factor EIF3 P35 Subunit and 60S Ribosomal protein L22 pseudogenes. Contains ESTs, RIKEN cDNA 2700038K18 gene, RIKEN cDNA 3110001N18 gene, expressed sequence AU041196,
1877	19268	NM_031104	q		ribosomal protein L22 EST, Weakly similar to RS9 RAT 40S RIBOSOMAL PROTEIN S9 [R.norvegicus], EST, Weakly similar to S55917 ribosomal protein S9, cytosolic [H.sapiens], Homo sapiens, clone IMAGE:4500773, mRNA, partial cds, RIKEN cDNA 3010033P07 gene, expressed sequence AL022771, expressed sequence AL022885, mitochondrial ribosomal protein S4,
1878	10878	NM_031108	q.bb		ribosomal protein S9 EST, Weakly similar to 40S RIBOSOMAL PROTEIN S11 [R.norvegicus], Homo sapiens mRNA; cDNA DKFZp434A0326 (from clone DKFZp434A0326), Human DNA sequence from clone RP5-1060K6 on chromosome 20p12.1-13. Contains an RPS11 (40S ribosomal protein S11) pseudogene, ESTs, STSs and GSSs, RAD21 homolog (S. pombe), ribosomal protein S11

TABLE S	RUMANI 3	HOMOFOGAE VK	REMOTRATION		Any, Docket No. 44921-5039WO Doc. No. 1793397.1
899. ID.	refittación	GenBenk Ace./ Ref. Seq. ID No.	Model (906)	Homologovs Cane Name	Homologous Cluster Name
				33,351	EST, Moderately similar to 40S RIBOSOMAL PROTEIN S21 [R.norvegicus], Human DNA sequence from clone RP5-1116H23 on chromosome 20 Contains a novel gene, a 40S ribosomal protein S21 pseudogene, 2 CpG islands, ESTs, STSs and GSSs, RIKEN cDNA 1810049N11 gene, RIKEN cDNA
1880	19162	NM_031111	aa		2410030A14 gene, ribosomal protein S21
					EST, Moderately similar to 40S RIBOSOMAL PROTEIN S21 [R.norvegicus], Human DNA sequence from clone RP5-1116H23 on chromosome 20 Contains a novel gene, a 40S ribosomal protein S21 pseudogene, 2 CpG islands, ESTs, STSs and GSSs, RIKEN cDNA 1810049N11 gene, RIKEN cDNA 2410030A14 gene, ribosomal protein
1880	19161	NM_031111	a,bb		S21 EST, Highly similar to 40S
					RIBOSOMAL PROTEIN S24 [R.norvegicus], EST, Weakly similar to JH0213 ribosomal protein S24, cytosolic [H.sapiens], EST, Weakly similar to RS24_HUMAN 40S RIBOSOMAL PROTEIN S24 [M.musculus], ESTs, Highly similar to JH0213 ribosomal protein S24, cytosolic [H.sapiens], ESTs, Weakly similar to RS24_HUMAN 40S RIBOSOMAL PROTEIN S24
1881	24615	NM_031112	a,y		[H.sapiens], ribosomal protein S24 EST, Moderately similar to S12583
					polyubiquitin 4 - mouse [M.musculus], EST, Moderately similar to ubiquitin / ribosomal protein S27a [R.norvegicus], EST, Weakly similar to R27A_HUMAN 40S RIBOSOMAL PROTEIN S27A (SUB 77-156 [H.sapiens], Homo sapiens cDNA FLJ11603 fis, clone HEMBA1003926, RIKEN cDNA 0610006J14 gene, expressed sequence Al132487, ribosomal protein
1882	20839	NM_031113	a,q		S27a, ubiquitin C EST, Moderately similar to A Chain A,
1883	19040	NM_031114	l,m,General	S100 calcium binding protein A10 (calgizzarin), S100 calcium-binding protein A10 (annexin II ligand, calpactin I, light polypeptide (p11))	P11 [H.sapiens], EST, Moderately similar to CALPACTIN I LIGHT CHAIN [R.norvegicus], S100 calcium-binding protein A10 (annexin II ligand, calpactin I, light polypeptide (p11)), calcium binding protein A11 (calgizzarin)
1884	16349	NM_031115	u		ESTs, Highly similar to 2111411A secretin receptor [H.sapiens], ESTs, Weakly similar to vasoactive intestinal polypeptide 1 [M.musculus], adenylate cyclase activating polypeptide 1 receptor 1, secretin receptor, vasoactive intestinal peptide receptor 1

TABLES	MANUH &	HOMOLOGNE AK	ROTATIONS		Atty, Doctor No. 44921-5039W0 Doc. No. 1793397.1
899. ID +	ldentifier	Centent Acel Ref. Seq. 10 No.	Model Gode	Hamalogous Gare	Homologous Gluster Name
					ESTs, Highly similar to SUOX RAT SULFITE OXIDASE PRECURSOR [R.norvegicus], RIKEN cDNA 0610009N12 gene, RIKEN cDNA 1810044O22 gene, RIKEN cDNA
1885	14970	NM_031127	General		2810034J18 gene, fatty acid desaturase 2, sulfite oxidase
				thyroid hormone receptor alpha, thyroid hormone receptor, alpha (avian erythroblastic leukemia viral (y-erb-a) oncogene	EST, Weakly similar to A30893 thyroid hormone receptor alpha, splice form 2 [H.sapiens], expressed sequence AW259572, expressed sequence R75201, thyroid hormone receptor, alpha (avian erythroblastic leukemia
1886	1814	NM_031134	n,q	homolog)	viral (v-erb-a) oncogene homolog) Kruppel-like factor 15 (kidney), Kruppel
1887	13359	NM 031135	General		like factor 9, RIKEN cDNA 4930480I16 gene, TGFB inducible early growth response, basic transcription element binding protein 1, trans-acting transcription factor 3, trans-acting transcription factor 6
1888	15052	NM_031136	а		ESTs, Highly similar to A38682 thymosin beta-4 [H.sapiens], ESTs, Highly similar to TYB4 MOUSE THYMOSIN BETA-4 [M.musculus], ESTs, Highly similar to TYB4_HUMAN THYMOSIN BETA-4 [H.sapiens], Human interferon-inducible mRNA (cDNA 6-26), expressed sequence AW544206, thymosin, beta 10, thymosin, beta 4, X chromosome, thymosin, beta 4, Y chromosome
1888	19359	NM_031136	а		
					EST, Moderately similar to A25074 vimentin [H.sapiens], EST, Weakly similar to A25074 vimentin [H.sapiens], ESTs, Moderately similar to VIME RAT VIMENTIN [R.norvegicus], ESTs, Weakly similar to A25074 vimentin [H.sapiens], ESTs, Weakly similar to VIME RAT VIMENTIN [R.norvegicus],
1889	15185	NM_031140	General		vimentin EST, Weakly similar to ACTB HUMAN
					ACTIN, CYTOPLASMIC 1 [R.norvegicus], ESTs, Highly similar to ATHUB actin beta [H.sapiens], ESTs, Weakly similar to ACTB_HUMAN ACTIN, CYTOPLASMIC 1 [R.norvegicus], Homo sapiens FKSG30 (FKSG30) mRNA, complete cds, RIKEN cDNA 1700052K15 gene, RIKEN cDNA 1700061J02 gene, actin, beta, actin-like 7a, actin-related protein
1890	21625	NM_031144	а,е		3-beta, melanoma X-actin CATX-8 protein, ESTs, Weakly similar
1004	200	NNA 024450			to R11A_HUMAN RAS-RELATED PROTEIN RAB-11A [R.norvegicus], RAB, member of RAS oncogene family like 2A, RAB11A, member RAS oncogene family, RAB11a, member RAS oncogene family, RAB25, member RAS oncogene family, RIKEN
1891	238	NM_031152	bb	<u> </u>	cDNA 2700023P08 gene

TABLES	HUMAN	HOMOLOGUE AN	- EMOTIVATIONI		Any, Docket No. 44921-5039W0 Doc. No. 1793397.1
800, ID No.	ldentifier	ConBank Ace.// Ref. Seq. ID No.	Model Code	Hemelogous Cene Name	Homologous Cluster Name
1891	240	NM 031152	bb		CATX-8 protein, ESTs, Weakly similar to R11A_HUMAN RAS-RELATED PROTEIN RAB-11A [R.norvegicus], RAB, member of RAS oncogene family. like 2A, RAB11A, member RAS oncogene family, RAB11a, member RAS oncogene family, RAB25, member RAS oncogene family, RAB25, CDNA 2700023P08 gene
1892	15277	NM 031237			EST, Moderately similar to UB5B_HUMAN UBIQUITIN-CONJUGATING ENZYME E2-17 KD 2 [R.norvegicus], ESTs, Moderately similar to 159365 ubiquitin conjugating enzyme [H.sapiens], ESTs, Moderately similar to UB5B_HUMAN UBIQUITIN-CONJUGATING ENZYME E2-17 KD 2 [M.musculus], RIKEN cDNA 1100001F19 gene, RIKEN cDNA 1600028117 gene, RIKEN cDNA 2700084L22 gene, Rattus norvegicus clone ubc2e ubiquitin conjugating enzyme (E217kB) mRNA, complete cds, expressed sequence AL022654, ubiquitin-conjugating enzyme E2D 1 (homologous to yeast UBC4/5), ubiquitin-conjugating enzyme E2D 2 (homologous to yeast UBC4/5), ubiquitin-conjugating enzyme E2D 3 (homologous to yeast UBC4/5), ubiquitin-conjugating enzyme E2D 3 (homologous to yeast UBC4/5)
1092	13211	14W_031237	g		ESTs, Weakly similar to YZ28_HUMAN HYPOTHETICAL PROTEIN ZAP128 [H.sapiens], Mus musculus, Similar to cytosolic acyl- CoA thioesterase 1, clone MGC:27572
1893	18083	NM_031315	q		IMAGE:4485973, mRNA, complete cds

TABLE 3	: HUMAN	HOMOLOGUEAK	ROTATIONS		Atty; Docket No. 44921-5039WO Doc. No. 1793397.1
Seq. ID No.	ldentifier	Cenbenk Acc./ Ref. Seq. ID No.	Model Code	Homologous Come Name	Hemologous Civstet Name
					PTE2_HUMAN PEROXISOMAL ACYL-COENZYME A THIOESTER HYDROLASE 2 (PEROXISOMAL LONG-CHAIN ACYL-COA THIOESTERASE 2) (ZAP128) [H.sapiens], ESTs, Moderately similar to JE0267 long-chain fatty-acyl-CoA hydrolase (EC 3.1.2) peroxisome proliferator-inducible - rat [R.norvegicus], ESTs, Moderately similar to PTE2_HUMAN PEROXISOMAL ACYL-COENZYME A THIOESTER HYDROLASE 2 (PEROXISOMAL LONG-CHAIN ACYL- COA THIOESTERASE 2) (ZAP128) [H.sapiens], ESTs, Weakly similar to PTE2_HUMAN PEROXISOMAL ACYL- COENZYME A THIOESTER HYDROLASE 2 (PEROXISOMAL LONG-CHAIN ACYL-COA THIOESTERASE 2) (ZAP128) [H.sapiens], ESTs, Weakly similar to YZ28_HUMAN HYPOTHETICAL PROTEIN ZAP128 [H.sapiens], Mus musculus, Similar to cytosolic acyl- COA thioesterase 1, clone MGC:27572 IMAGE:4485973, mRNA, complete
				cytosolic acyl-CoA thioesterase 1,	cds, RIKEN cDNA 4632408A20 gene, cytosolic acyl-CoA thioesterase 1,
1893	1858	NM_031315	q	peroxisomal long-chain acyl-coA thioesterase	expressed sequence AW108394, peroxisomal long-chain acyl-coA It-complex testis expressed 1, t-
1894	15663	NM_031318	General		complex-associated-testis-expressed 1 like 1
1895	1422	NM 031324	bb,General		ESTs, Moderately similar to I38134 prolyl oligopeptidase [H.sapiens], prolyl endopeptidase
1896	18597	NM 031325	g,bb		UDP-glucose dehydrogenase
1897	11259	NM_031327	i,cc,General		ESTs, Moderately similar to CYR6 MOUSE CYR61 PROTEIN PRECURSOR [M.musculus], cysteine rich protein 61, cysteine-rich, angiogenic inducer, 61
1898	4235	NM_031330	General	heterogeneous nuclear ribonucleoprotein A/B	ESTs, Highly similar to WZHURS argininosuccinate lyase [H.sapiens], ESTs, Weakly similar to 1601424A argininosuccinate lyase [R.norvegicus], Homo sapiens cDNA FLJ14312 fis, clone PLACE3000322, Musashi-1 homolog (Drosophila), RIKEN cDNA 2510006M18 gene, RIKEN cDNA 4933434H11 gene, argininosuccinate lyase, heterogeneous nuclear ribonucleoprotein A/B, heterogeneous nuclear ribonucleoprotein D-like EST, Weakly similar to PSD4_HUMAN 26S PROTEASOME REGULATORY SUBUNIT S5A [H.sapiens], ESTs, Moderately similar to PSD4_HUMAN 26S PROTEASOME REGULATORY
1899	18375	NM_031331	l,m		SUBUNIT S5A [H.sapiens], proteasome (prosome, macropain) 26S subunit, non-ATPase, 4

TABLE		HOMOTOGAL	SHOITATON		. (Alty: Docket No. 44921-5039WC Doc. No. 1793397.1
Seq. (D) No. ::	Identifier	ConBank Ace./ Ref. Seq. ID No.	Model Code	Hamalagaus Cena	Homologous Cluster Name
1900	3519	NM_031334	cc	cadherin 1, cadherin 1, type 1, E-cadherin (epithelial)	ESTs, Weakly similar to I49556 cadherin-11 - mouse [M.musculus], RIKEN cDNA 2610005L07 gene, cadherin 1, type 1, E-cadherin (epithelial), cadherin 6, cadherin 6, type 2, K-cadherin (fetal kidney)
1901	20698	NM 031357	b		
1903	634	NM_031509	n		EST, Moderately similar to GTC MOUSE GLUTATHIONE S- TRANSFERASE YC [M.musculus], glutathione S-transferase A3, glutathione S-transferase, alpha 3
1903	25525	NM_031509	n		
1903	25069	NM_031509	b,n,w		
1903	635	NM_031509	z		EST, Moderately similar to GTC MOUSE GLUTATHIONE S- TRANSFERASE YC [M.musculus], glutathione S-transferase A3, glutathione S-transferase, alpha 3 EST, Highly similar to RON HUMAN
				met proto-oncogene, met proto-oncogene (hepatocyte growth factor	MACROPHAGE-STIMULATING PROTEIN RECEPTOR PRECURSOR [H.sapiens], ESTs, Highly similar to TVHUME hepatocyte growth factor receptor precursor [H.sapiens], Mus musculus D86 mRNA, complete cds, Rattus norvegicus ryk mRNA for tyrosine kinase-related protein, partial cds, macrophage stimulating 1 receptor (c-met-related tyrosine kinase), met proto-oncogene, met proto-oncogene (hepatocyte growth
1904	848	NM_031517		receptor)	factor receptor) RIKEN cDNA 0610007D04 gene, kallikrein 1, renal/pancreas/salivary, kallikrein 5, kallikrein 9, nerve growth factor, alpha, nerve growth factor,
1905	1872	NM 031523	а		gamma
1905	16245	NM_031523	a,d,u		EST, Moderately similar to epidermal growth factor binding protein type 1 [M.musculus], EST, Weakly similar to pre-pro-protein for kallikrein [H.sapiens] EST, Moderately similar to epidermal growth factor binding protein type 1 [M.musculus], EST, Weakly similar to pre-pro-protein for kallikrein
1005	16244	NM 031523	ا		
1905	16244	NM_031523	a	protein phosphatase 1, catalytic subunit, alpha	[H.sapiens] EST, Weakly similar to JN0723 phosphoprotein phosphatase [H.sapiens], protein phosphatase 1,
1906	9370	NM_031527	W General	small inducible cytokine A2, small inducible cytokine A2 (monocyte chemotactic protein 1, homologous to mouse Sig- je)	catalytic subunit, alpha isoform EST, Weakly similar to SY02 RAT SMALL INDUCIBLE CYTOKINE A2 PRECURSOR [R.norvegicus], expressed sequence Al323594, expressed sequence AW987545, small inducible cytokine A2, small inducible cytokine A24, small inducible cytokine A7 (monocyte chemotactic protein 3), small inducible cytokine subfamily A ((Cys-Cys), member 17

TABLE 8		HOMOLOGUE AN	ROTATIONS		Aiiy, Dockei No. 44921-5039WO Doc. No. 1793397.1
800. ID No	ldentifier	Confork Aced Ref. Seq. ID No.	Model Code	Homologous Gane) Mana	t Homologous Gluster Name
				small inducible cytokine A2, small inducible cytokine A2 (monocyte chemotactic protein 1, homologous to mouse Sig-	EST, Weakly similar to SY02 RAT SMALL INDUCIBLE CYTOKINE A2 PRECURSOR [R.norvegicus], expressed sequence Al323594, expressed sequence AW987545, small inducible cytokine A2, small inducible cytokine A24, small inducible cytokine A7 (monocyte chemotactic protein 3), small inducible cytokine subfamily A
1907	20449	NM_031530	General	je)	(Cys-Cys), member 17
					ESTs, Moderately similar to UDP-GLUCURONOSYLTRANSFERASE 2B5 PRECURSOR, MICROSOMAL [M.musculus], ESTs, Weakly similar to UDB7_HUMAN UDP-GLUCURONOSYLTRANSFERASE 2B7 PRECURSOR, MICROSOMAL [H.sapiens], ESTs, Weakly similar to UDBH_HUMAN UDP-GLUCURONOSYLTRANSFERASE 2B17 PRECURSOR, MICROSOMAL [H.sapiens], RIKEN cDNA 0610033E06 gene, UDP glycosyltransferase 2 family, polypeptide B17, UDP-glucuronosyltransferase 2 family, member 5, expressed sequence
1908	14633	NM_031533	u		AA986709
1909	16048	NM 031541	f	CD36 antigen (collagen type I receptor, thrombospondin receptor)- like 1, scavenger receptor class B1	CD36 antigen (collagen type I receptor, thrombospondin receptor)-like 1, CD36 antigen (collagen type I receptor, thrombospondin receptor)-like 2, ESTs, Weakly similar to JC5533 scavenger receptor class B type I precursor - rat [R.norvegicus], Homo sapiens scavenger receptor class B type III SR-BIII mRNA, partial cds, scavenger receptor class B1
1910	4011	NM_031543	c,q	cytochrome P450, 2e1, ethanol inducible, cytochrome P450, subfamily IIE (ethanol- inducible)	cytochrome P450, 2e1, ethanol inducible, cytochrome P450, subfamily IIE (ethanol-inducible)
1910	4010	NM_031543	c,q	cytochrome P450, 2e1, ethanol inducible, cytochrome P450, subfamily IIE (ethanol- inducible)	cytochrome P450, 2e1, ethanol inducible, cytochrome P450, subfamily IIE (ethanol-inducible)
1910	4012	NM_031543	q	cytochrome P450, 2e1, ethanol inducible, cytochrome P450, subfamily IIE (ethanol- inducible)	cytochrome P450, 2e1, ethanol inducible, cytochrome P450, subfamily IIE (ethanol-inducible)
1911	28	NM_031546	General	regucalcin, regucalcin (senescence marker protein-30)	regucalcin, regucalcin (senescence marker protein-30)
1912	24640	NM_031548	h,cc	sodium channel, nonvoltage-gated 1 alpha, sodium channel, nonvoltage-gated, type I, alpha polypeptide	expressed sequence AW742291, sodium channel, nonvoltage-gated 1 alpha, sodium channel, nonvoltage-gated 1, delta, sodium channel, nonvoltage-gated, type I, alpha polypeptide
1913	17149	NM_031549	x	transgelin	transgelin
1913	17151	NM_031549	х	transgelin	transgelin

TABLES	E HOMMAN	HOMOFOGAE VI	SKONATOKE		Aiiy. Dooker No. 44921-5039Wd Doc. No. 1793397.1
899. ID No.	ldentilier	CenBank Acc./ Ref. Seq. ID No.	Model Gode	Homologous Como Name	Homologous Auster Name
1914	13105	NM_031552	w	adducin 3 (gamma)	ESTs, Moderately similar to ADDG_MOUSE GAMMA ADDUCIN (ADDUCIN-LIKE PROTEIN 70) [M.musculus], adducin 3 (gamma) ESTs, Weakly similar to CPT1 MOUSE CARNITINE O-
				carnitine palmitoyltransferase 1,	PALMITOYLTRANSFERASE I, MITOCHONDRIAL LIVER ISOFORM [M.musculus], ESTs, Weakly similar to I59351 carnitine O-palmitoyltransferase [H.sapiens], ESTs, Weakly similar to MITOCHONDRIAL CARNITINE O-PALMITOYLTRANSFERASE I, LIVER ISOFORM [R.norvegicus], carnitine palmitoyltransferase 1, liver, carnitine
1915	15411	NM 031559	d,r	liver, carnitine palmitoyltransferase I, liver	palmitoyltransferase 1, muscle, carnitine palmitoyltransferase I, liver
1915	15411	14141_031333	u,i	Y box protein 1, nuclease sensitive element binding	ESTs, Highly similar to I39382 Y box- binding protein 1 - human [H.sapiens],
1916	16164	NM_031563	a,y	protein 1	RIKEN cDNA 1700102N10 gene EST. Moderately similar to 40S
					RIBOSOMAL PROTEIN S7 [R.norvegicus], EST, Weakly similar to 40S RIBOSOMAL PROTEIN S7 [R.norvegicus], EST, Weakly similar to
					RS7_HUMAN 40S RIBOSOMAL PROTEIN S7 [M.musculus], ESTs, Highly similar to JC4388 ribosomal protein S7, cytosolic [H.sapiens], ESTs, Highly similar to RS7_HUMAN 40S RIBOSOMAL PROTEIN S7
		NIA 004570		discount protein S7	[H.sapiens], nuclear factor of kappa light polypeptide gene enhancer in B- cells inhibitor-like 2, ribosomal protein
1917	9621	NM_031570	bb	ribosomal protein S7	S7 EST, Moderately similar to 40S RIBOSOMAL PROTEIN S7
					[R.norvegicus], EST, Weakly similar to 40S RIBOSOMAL PROTEIN S7 [R.norvegicus], EST, Weakly similar to
					RS7_HUMAN 40S RIBOSOMAL PROTEIN S7 [M.musculus], ESTs,
e E					Highly similar to JC4388 ribosomal protein S7, cytosolic [H.sapiens], ESTs, Highly similar to RS7_HUMAN
					40S RIBOSOMAL PROTEIN S7 [H.sapiens], nuclear factor of kappa light polypeptide gene enhancer in B-
1917	9620	NM 031570	w,bb	ribosomal protein S7	cells inhibitor-like 2, ribosomal protein
1311	5520	001070	11,00		ESTs, Moderately similar to KPBG_HUMAN PHOSPHORYLASE B
					KINASE GAMMA CATALYTIC CHAIN, SKELETAL MUSCLE ISOFORM [H.sapiens], ESTs, Moderately similar
					to PHOSPHORYLASE B KINASE GAMMA CATALYTIC CHAIN, SKELETAL MUSCLE ISOFORM
					[R.norvegicus], RIKEN cDNA 1500017l02 gene, endoplasmic reticulum (ER) to nucleus signalling 2,
1918	546	NM_031573	f	phosphorylase kinase gamma, phosphorylase kinase, gamma 1 (muscle)	phosphorylase kinase gamma, phosphorylase kinase, gamma 1 (muscle)

Good and			INOTATIONS		Affy. Docket No. 44921-5039W0 Doc. No. 1798897.1
Seg. ID (No.	ldendiler	ConBank Ace <i>l</i> Ref. Seq. ID No.	Modal Goda	Homologous Como Namo	Homologous Cluster Name
					ESTs, Highly similar to A Chain A, Crystal Structure Of The Fmn-Binding Domain Of Human Cytochrome P450 Reductase At 1.93a Resolution {SUB 61-241 [H.sapiens], NADPH- dependent FMN and FAD containing
				P450 (cytochrome)	oxidoreductase, P450 (cytochrome) oxidoreductase, RIKEN cDNA 4930447P04 gene, hypothetical
1919	1921	NM_031576	f	oxidoreductase	protein FLJ10900 ESTs, Highly similar to A Chain A,
				DATE (s. d. charana)	Crystal Structure Of The Fmn-Binding Domain Of Human Cytochrome P450 Reductase At 1.93a Resolution (SUB 61-241 [H.sapiens], NADPH- dependent FMN and FAD containing oxidoreductase, P450 (cytochrome) oxidoreductase, RIKEN cDNA
1919	1920	NM_031576	r	P450 (cytochrome) oxidoreductase	4930447P04 gene, hypothetical protein FLJ10900
1920	24219	NM_031579	i,General	protein tyrosine phosphatase 4a1, protein tyrosine phosphatase type IVA, member 1	protein tyrosine phosphatase 4a1, protein tyrosine phosphatase 4a2, protein tyrosine phosphatase type IVA, member 1, protein tyrosine phosphatase type IVA, member 2, protein tyrosine phosphatase type IVA, member 3
				solute carrier family 22 (organic cation	EST, Moderately similar to JC4884 organic cation transporter protein 2 - rat [R.norvegicus], EST, Weakly similar to OCN2 MOUSE ORGANIC CATION/CARNITINE TRANSPORTER 2 [M.musculus], ESTs, Highly similar to OCN2 HUMAN ORGANIC CATION/CARNITINE TRANSPORTER 2 [H.sapiens], ESTs, Highly similar to organic cation transporter [H.sapiens], organic cationic transporter-like 1, solute carrier family 22 (organic cation transporter), member 1, solute carrier family 22 (organic cation transporter), member 4, solute carrier family 22 (organic cation transporter), member 5, solute carrier family 22 (organic cation transporter), member 5, solute carrier family 22 (organic cation transporter), member 9, solute
1921	770	NM_031584	k,x	transporter), member 2	carrier family 22, member 3 ESTs, Highly similar to NRG2_MOUSE
					PRO-NEUREGULIN-2 PRECURSOR (PRO-NEUREGULIN-2 PRECURSOR (PRO-NRG2) [CONTAINS: NEUREGULIN-2 (NRG-2) (DIVERGENT OF NEUREGULIN 1) (DON-1)] [M.musculus], ESTs, Weakly similar to NRG2_MOUSE PRO-NEUREGULIN-2 PRECURSOR (PRO-NRG2) [CONTAINS: NEUREGULIN-2 (NRG-2) (DIVERGENT OF NEUREGULIN 1) (DON-1)]

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TABLES	K WWWH I	HOMOROGALE W	EKIOTTATOKI		Any. Docket No. 44921-51119WO Doc. No. 1793397.1
Scg. [D] No.	ldenillier	Combank Acel Ref. Seq. ID No.	Model Gode	Homologous Gane Mandalla and Mandalla logous Cluster Name	
					ESTs, Highly similar to NRG2_MOUSE PRO-NEUREGULIN-2 PRECURSOR (PRO-NRG2) [CONTAINS: NEUREGULIN-2 (NRG-2) (DIVERGENT OF NEUREGULIN 1) (DON-1)] [M.musculus], ESTs, Weakly similar to NRG2_MOUSE PRO- NEUREGULIN-2 PRECURSOR (PRO-
.0.					NRG2) [CONTAINS: NEUREGULIN-2 (NRG-2) (DIVERGENT OF NEUREGULIN 1) (DON-1)]
1922	18005	NM_031588	h		[M.musculus], neuregulin 1 ESTs, Highly similar to NRG2_MOUSE PRO-NEUREGULIN-2 PRECURSOR (PRO-NRG2) [CONTAINS: NEUREGULIN-2 (NRG-2) (DIVERGENT OF NEUREGULIN 1) (DON-1)] [M.musculus], ESTs, Weakly similar to NRG2_MOUSE PRO- NEUREGULIN-2 PRECURSOR (PRO- NRG2) [CONTAINS: NEUREGULIN-2 (NRG-2) (DIVERGENT OF
1922	18011	NM_031588	cc,General	N-0-	NEUREGULIN 1) (DON-1)] [M.musculus], neuregulin 1 EST, Moderately similar to PRSA RAT
4000		NM 024505		proteasome (prosome, macropain) 26S subunit, ATPase 3, proteasome (prosome, macropain) 26S	26S PROTEASE REGULATORY SUBUNIT 6A [R.norvegicus], EST, Weakly similar to PRS4 MOUSE 26S PROTEASE REGULATORY SUBUNIT 4 [M.musculus], EST, Weakly similar to PRSA RAT 26S PROTEASE REGULATORY SUBUNIT 6A [R.norvegicus], ESTs, Moderately similar to PRSA RAT 26S PROTEASE REGULATORY SUBUNIT 6A [R.norvegicus], expressed sequence Al325227, protease (prosome, macropain) 26S subunit, ATPase 1, proteasome (prosome, macropain) 26S subunit, ATPase 3, proteasome (prosome, macropain) 26S subunit,
1923	1584	NM_031595	k	subunit, ATPase, 3	ATPase, 3 Mus musculus adult male small
1924	24235	NM 031614	lv	. thioredoxin reductase 1	intestine cDNA, RIKEN full-length enriched library, clone:2010001F03, full insert sequence, glutathione reductase 1, thioredoxin reductase 1, thioredoxin reductase 2, thioredoxin reductase beta
1924	24234	NM_031614	General	thioredoxin reductase 1	Mus musculus adult male small intestine cDNA, RIKEN full-length enriched library, clone:2010001F03, full insert sequence, glutathione reductase 1, thioredoxin reductase 1, thioredoxin reductase 2, thioredoxin reductase beta
1925	1639	NM_031627	j.l,v	nuclear receptor subfamily 1, group H, member 3	EST, Moderately similar to A56043 steroid hormone receptor-like protein RLD-1 - rat [R.norvegicus], expressed sequence AU018371, nuclear receptor subfamily 1, group H, member 3, nuclear receptor subfamily 1, group H, member 4

TABLE		H <mark>OWOTOGNE</mark> W			Atty. Docket No. 44921-5039W0 Doc. No. 1793397.1
Sog. ID No.	ldenilliter	ConBank Aced Ref. Seq. (D No.	Model Gode 🦠	Homologous Ceme Name	Homologous Gluster Name
					EST, Moderately similar to CPBP RAT CORE PROMOTER ELEMENT-BINDING PROTEIN [R.norvegicus], ESTs, Moderately similar to CPBP RAT CORE PROMOTER ELEMENT-BINDING PROTEIN [R.norvegicus], Kruppel-like factor 3 (basic), Kruppellike factor 7
1926	1727	NM_031642	m,General	core promoter element binding protein	(ubiquitous), core promoter element binding protein
				mitogen activated protein kinase kinase 2, mitogen- activated protein kinase	ESTs, Highly similar to MPK1 MOUSE DUAL SPECIFICITY MITOGEN-ACTIVATED PROTEIN KINASE KINASE 1 [M.musculus], ESTs, Moderately similar to MPK1_HUMAN DUAL SPECIFICITY MITOGEN-ACTIVATED PROTEIN KINASE KINASE 1 [H.sapiens], Mus musculus 12 days embryo head cDNA, RIKEN full-length enriched library, clone:3000002B10, full insert sequence, mitogen activated protein kinase kinase 1, mitogen activated protein kinase kinase 3, mitogen activated protein kinase kinase 7, mitogen-activated protein kinase kinase 1, mitogen-activated protein kinase kinase 1, mitogen-activated protein kinase kinase 1, mitogen-activated protein kinase
1927	20766	NM_031643	y	kinase 2	kinase kinase 7
					latexin, latexin protein, retinoic acid receptor responder (tazarotene
1929	1993	NM_031655	k,l,m,General	latexin, latexin protein	induced) 1 Human DNA sequence from clone RP5-822J19 on chromosome 20. Contains an alpha-endosulfine pseudogene, STSs and GSSs, cyclic AMP phosphoprotein, 19 kD, cyclic AMP phosphoprotein, 19kD,
1930	2057	NM_031660	e	solute carrier family 15 (H+/peptide transporter),	endosulfine alpha EST, Moderately similar to PET2 RAT OLIGOPEPTIDE TRANSPORTER, KIDNEY ISOFORM [R.norvegicus], EST, Moderately similar to PET2_HUMAN OLIGOPEPTIDE TRANSPORTER, KIDNEY ISOFORM [H.sapiens], expressed sequence C78862, solute carrier family 15
1931	15039	NM_031672	k,General	member 2	(H+/peptide transporter), member 2 2-4-dienoyl-Coenzyme A reductase 2, peroxisomal, ESTs, Weakly similar to HCD2 RAT 3-HYDROXYACYL-COA DEHYDROGENASE TYPE II [R.norvegicus], H2-K region expressed gene 6, hydroxyacyl-Coenzyme A dehydrogenase, type II, hydroxyprostaglandin dehydrogenase 15 (NAD), hydroxysteroid (17-beta) dehydrogenase 10, hypothetical protein FLJ14431, retinal short-chain
1932	15175	NM_031682	bb	golgi SNAP receptor	dehydrogenase/reductase retSDR3 golgi SNAP receptor complex member
1933	1004	NM_031685	v	complex member 2	2

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				75 Sale Line Brasiling	<u>Doc. No. 1793397.1</u>
(10) (10)	letantifer	ConBank Ace./ Ref. Seq. ID No.	Model Code	Homologous Cene Name	Homologous Cluster Name
1934	19727	NM 031687	a,q,s	ubiquitin A-52 residue ribosomal protein fusion product 1	EST, Moderately similar to I65237 ubiquitin/ribosomal protein L40 - rat [R.norvegicus], Homo sapiens ubiquitin-like fusion protein mRNA, complete cds, RIKEN cDNA 0610006J14 gene, Rattus norvegicus RSD-7 mRNA, complete cds, ubiquitin A-52 residue ribosomal protein fusion product 1
					ESTs, Weakly similar to A39484 androgen-withdrawal apoptosis protein RVP1, prostatic - rat [R.norvegicus], claudin 12, claudin 3, expressed
1935	20404	NM_031700	j,r,y	claudin 3	sequence Al182374 ESTs, Weakly similar to A39484
1935	20405	NM_031700	o,r	claudin 3	androgen-withdrawal apoptosis protein RVP1, prostatic - rat [R.norvegicus], claudin 12, claudin 3, expressed sequence Al182374
1936	811	NM_031705	General	dihydropyrimidinase	PRO0195 protein, collapsin response mediator protein 5, collapsin response mediator protein-5; CRMP3-associated molecule, dihydropyrimidinase, dihydropyrimidinase-like 2, dihydropyrimidinase-related protein
1936	812	NM_031705	o,v,bb,General	dihydropyrimidinase	PRO0195 protein, collapsin response mediator protein 5, collapsin response mediator protein-5; CRMP3-associated molecule, dihydropyrimidinase, dihydropyrimidinase-like 2, dihydropyrimidinase-related protein EST, Moderately similar to
1937	16204	NM_031706	q,bb		RS8_HUMAN 40S RIBOSOMAL PROTEIN S [H.sapiens], EST, Weakly similar to 40S RIBOSOMAL PROTEIN S8 [M.musculus], EST, Weakly similar to 40S RIBOSOMAL PROTEIN S8 [R.norvegicus], ESTs, Moderately similar to RS8_HUMAN 40S RIBOSOMAL PROTEIN S [H.sapiens], RIKEN cDNA 1110008P08 gene, ribosomal protein S8
					EST, Moderately similar to RS8_HUMAN 40S RIBOSOMAL PROTEIN S [H.sapiens], EST, Weakly similar to 40S RIBOSOMAL PROTEIN S8 [M.musculus], EST, Weakly similar to 40S RIBOSOMAL PROTEIN S8 [R.norvegicus], ESTs, Moderately similar to RS8_HUMAN 40S RIBOSOMAL PROTEIN S [H.sapiens], RIKEN cDNA 1110008P08 gene,
1937	16205	NM_031706	a,y		ribosomal protein S8 ESTs, Weakly similar to G100_HUMAN 110 KDA CELL MEMBRANE GLYCOPROTEIN
					[H.sapiens], cell membrane
1938	24081	NM_031708	m		glycoprotein, 110000M(r) (surface antigen) ESTs, Highly similar to R3HU12
					ribosomal protein S12, cytosolic [H.sapiens], mitochondrial ribosomal
1939	16918	NM_031709	a,q	<u> </u>	protein L50, ribosomal protein S12

TABLE 8	HUMAN	HOMOFOGAE VIX	EKIÖ ITATOKI		Ally, Dockel No. 44921-5039WO Doc. No. 1793997:1
Seq. ID: No III:		Cenbank Ace./ Ref. Seq. ID No.	Model Gode	Homologous Cone Name	Homologous Elvster Namo
					ESTs, Weakly similar to T30259 multiple PDZ domain protein - mouse [M.musculus], ESTs, Weakly similar to T46612 multi PDZ domain protein 1 - rat [R.norvegicus], PDZ domain containing 1, channel-interacting PDZ domain protein, hypothetical protein FLJ22756, multiple PDZ domain protein, semaF cytoplasmic domain associated protein 3, solute carrier family 9 (sodium/hydrogen exchanger)
					isoform 3 regulatory factor 1, solute carrier family 9 (sodium/hydrogen exchanger), isoform 3 regulatory factor
1940	1081	NM_031712	General	PDZ domain containing 1	2, syntaxin binding protein 4 ESTs, Highly similar to S71429
					phosphofructokinase, muscle - rat [R.norvegicus], Mus musculus adult male stomach cDNA, RIKEN full-length enriched library, clone:2210403E17, full insert sequence, expressed
1941	1340	NM_031715	b,n,u,cc,General	phosphofructokinase, muscle	sequence AA407869, phosphofructokinase, liver, B-type, phosphofructokinase, muscle
				family, member A2, aldehyde dehydrogenase	ESTs, Weakly similar to DHA4 RAT FATTY ALDEHYDE DEHYDROGENASE [R.norvegicus], RIKEN cDNA 1700001N19 gene, RIKEN cDNA 1700055N04 gene, aldehyde dehydrogenase 3 family, member A2, aldehyde dehydrogenase family 3, subfamily A2, expressed
1942	23884	NM_031731	j,s	family 3, subfamily A2	sequence Al848594
		÷			ESTs, Highly similar to N-ACETYLLACTOSAMINE SYNTHASE [M.musculus], UDP-Gal:betaGlcNAc beta 1,4- galactosyltransferase, polypeptide 1, UDP-Gal:betaGlcNAc beta 1,4- galactosyltransferase, polypeptide 2, UDP-Gal:betaGlcNAc beta 1,4- galactosyltransferase, polypeptide 3, UDP-Gal:betaGlcNAc beta 1,4- galactosyltransferase, polypeptide 4, UDP-Gal:betaGlcNAc
1943	10241	NM 031740	d	UDP-Gal:betaGlcNAc beta 1,4- galactosyltransferase, polypeptide 6, UDP- Gal:betaGlcNAc beta 1,4- galactosyltransferase, polypeptide 6	beta 1,4- galactosyltransferase, polypeptide 5, UDP-Gal:betaGlcNAc beta 1,4- galactosyltransferase, polypeptide 6, UDP-Gal:betaGlcNAc beta 1,4-galactosyltransferase, polypeptide 3
1944	1214	NM 031741	ır	solute carrier family 2 (facilitated glucose transporter), member 5, solute carrier family 2 (facilitated glucose/fructose transporter), member 5	Mus musculus, clone MGC:8298 IMAGE:3593581, mRNA, complete cds, glucose transporter protein 11, solute carrier family 2 (facilitated glucose transporter), member 5, solute carrier family 2 (facilitated glucose transporter), member 9, solute carrier family 2 (facilitated glucose/fructose transporter), member 5

TABLE 8	HUMANI	HOMOLOGUE AN	ROTATIONS		Aity, Docket No. 44921-5089W0 Doc. No. 1798897.1
Seq. 10 No.	(dentifier	Cenbank Ace <i>l</i> Ref. Seq. ID No.	Model Gode 🕠	Hemologous Game Name	Homologous Gluster Namo
1944	1215	NM_031741	r	solute carrier family 2 (facilitated glucose transporter), member 5, solute carrier family 2 (facilitated glucose/fructose transporter), member 5	Mus musculus, clone MGC:8298 IMAGE:3593581, mRNA, complete cds, glucose transporter protein 11, solute carrier family 2 (facilitated glucose transporter), member 5, solute carrier family 2 (facilitated glucose transporter), member 9, solute carrier family 2 (facilitated glucose/fructose transporter), member 5 ESTs, Highly similar to C166_HUMAN
1945	20724	NM_031753	h		CD166 ANTIGEN PRECURSOR [H.sapiens], Lutheran blood group (Auberger b antigen included), activated leucocyte cell adhesion molecule, activated leukocyte cell adhesion molecule, advanced glycosylation end product-specific receptor, melanoma cell adhesion molecule
				,	EST, Weakly similar to LIS1 MOUSE PLATELET-ACTIVATING FACTOR ACETYLHYDROLASE IB ALPHA SUBUNIT [R.norvegicus], ESTs, Weakly similar to LIS1 MOUSE PLATELET-ACTIVATING FACTOR ACETYLHYDROLASE IB ALPHA SUBUNIT [R.norvegicus], ESTs, Weakly similar to LIS1_HUMAN PLATELET-ACTIVATING FACTOR ACETYLHYDROLASE IB ALPHA SUBUNIT [H.sapiens], F-box protein FBW7, KIAA0007 protein, U3 snoRNP-associated 55-kDa protein, f-box and WD-40 domain protein 2, platelet-activating factor acetylhydrolase, isoform 1b, beta1 subunit, platelet-activating factor acetylhydrolase,
1946	20753	NM_031763	h		isoform Ib, alpha subunit (45kD) EST, Weakly similar to LIS1 MOUSE PLATELET-ACTIVATING FACTOR ACETYLHYDROLASE IB ALPHA SUBUNIT [R.norvegicus], ESTs, Weakly similar to LIS1 MOUSE PLATELET-ACTIVATING FACTOR ACETYLHYDROLASE IB ALPHA SUBUNIT [R.norvegicus], ESTs, Weakly similar to LIS1_HUMAN PLATELET-ACTIVATING FACTOR ACETYLHYDROLASE IB ALPHA SUBUNIT [H.sapiens], F-box protein FBW7, KIAA0007 protein, U3 snoRNP- associated 55-kDa protein, F-box and WD-40 domain protein 2, platelet- activating factor acetylhydrolase, isoform 1b, beta1 subunit, platelet- activating factor acetylhydrolase,
1946	20752	NM_031763	у	Rab acceptor 1	isoform lb, alpha subunit (45kD)
1947 1948	14953 14184	NM_031774 NM_031776 NM_031776	t,General d,o,t,General	(prenylated)	guanine deaminase guanine deaminase

TABLES		HQWQTQGM3 (A)	SVOTATIONS		Aity. Docket No. 44921-5039W0 Doc. No. 1793397.1
Sog. (D No.	ldentlifter	GenBank Aced Ref. Seq. 10 No.	Model Gode	Homologous Cene Name	Homologous Gluster Namo
bet the first section in teams		V			ESTs, Highly similar to NFL2 RAT NUCLEAR FACTOR ERYTHROID 2 RELATED FACTOR 2 [R.norvegicus], ESTs, Weakly similar to NFL2 RAT NUCLEAR FACTOR ERYTHROID 2 RELATED FACTOR 2 [R.norvegicus], nuclear factor (erythroid-derived 2)-like
1949	1169	NM_031789	c		2, nuclear, factor, erythroid derived 2, like 2
1950	16155	NM_031810	d,z		defensin beta 1, defensin beta 2, defensin, beta 1
1950	16156	NM_031810	d		defensin beta 1, defensin beta 2, defensin, beta 1 EST, Weakly similar to T42627 ADP-
				G protein-coupled receptor	ribosylation factor-directed GTPase activating protein, isoform a - mouse [M.musculus], ESTs, Highly similar to T42627 ADP-ribosylation factor-directed GTPase activating protein, isoform a - mouse [M.musculus], G protein-coupled receptor kinase-interactor 1, G protein-coupled receptor kinase-interactor 2, Homo sapiens p95 paxillin-kinase linker mRNA, complete cds, RIKEN cDNA 1700030C10 gene, development and
1951	17194	NM_031814	z	kinase-interactor 1	differentiation enhancing ESTs, Weakly similar to
1952	17535	NM_031816	bb		GBLP_HUMAN GUANINE NUCLEOTIDE-BINDING PROTEIN BETA SUBUNIT-LIKE PROTEIN 12.3 [R.norvegicus], Homo sapiens cDNA: FLJ21913 fis, clone HEP03888, Homo sapiens, clone IMAGE:3502107, mRNA, partial cds, RAE1 (RNA export 1, S.pombe) homolog, WD repeat domain 10, expressed sequence Al173248, expressed sequence Al504353, guanine nucleotide binding protein (G protein), beta polypeptide 2- like 1, guanine nucleotide binding protein, beta 2, related sequence 1, retinoblastoma binding protein 7, retinoblastoma-binding protein 7, transducin (beta)-like 2
					ESTS, Highly similar to A57286 probable serine/threonine protein kinase [M.musculus], ESTS, Highly similar to SNK_RAT SERINE/THREONINE-PROTEIN KINASE SNK (SERUM INDUCIBLE KINASE) [R.norvegicus], ESTS, Weakly similar to SNK_RAT SERINE/THREONINE-PROTEIN KINASE SNK (SERUM INDUCIBLE KINASE) [R.norvegicus], cytokine-inducible kinase, serine/threonine kinase 18, serum-inducible kinase,
1953	2655	NM_031821	i,l,m,aa	serum-inducible kinase	tousled-like kinase 2 (Arabidopsis) flotillin 1, flotillin 2
1954	22321	NM_031830 NM_031832	o,t,u,General	lectin, galactose binding, soluble 3, lectin, galactoside-binding, soluble, 3 (galectin 3)	EST, Weakly similar to X-Ray Crystal Structure Of The Human Galectin-3 Carbohydrate Recognition Domain [H.sapiens], galectin-related inter-fiber protein

TABLE 8	E HUMAN	HOMOLOGUE AN	INOTATIONS	e produktor Albier (8 di	Aity. Docket No. 44921-5089W0 Doc. No. 1793397.1
Seg. (D No.	ldenililer	CenBenk Ace <i>l</i> Ref. Seq. 10 No.	Model Gode	Hemologous Cene Name	Momologovs Cluster Name
					expressed sequence Al266890, expressed sequence Al853643, sulfotransferase family 1A, phenol- preferring, member 1, sulfotransferase
					family 4A, member 1, sulfotransferase family, cytosolic, 1A, phenol-preferring, member 1, sulfotransferase family, cytosolic, 1A, phenol-preferring,
1956	4748	NM 031834	e,t		member 2, sulfotransferase family, cytosolic, 1A, phenol-preferring, member 3
					expressed sequence Al266890, expressed sequence Al853643, sulfotransferase family 1A, phenol- preferring, member 1, sulfotransferase
					family 4A, member 1, sulfotransferase family, cytosolic, 1A, phenol-preferring, member 1, sulfotransferase family, cytosolic, 1A, phenol-preferring,
1956	4749	NM 031834	le,t		member 2, sulfotransferase family, cytosolic, 1A, phenol-preferring, member 3
					ESTs, Weakly similar to AGT2 RAT ALANINEGLYOXYLATE AMINOTRANSFERASE 2 PRECURSOR [R.norvegicus], RIKEN
					cDNA 1300019H02 gene, RIKEN cDNA 2900006B13 gene, alanine- glyoxylate aminotransferase 2, alanine- glyoxylate aminotransferase 2-like 1,
1957	7914	NM_031835	е		ornithine aminotransferase c-fos induced growth factor (vascular endothelial growth factor D), vascular
1958	8385	NM_031836	h		endothelial growth factor, vascular endothelial growth factor B c-fos induced growth factor (vascular
1958	8384	NM_031836	h		endothelial growth factor D), vascular endothelial growth factor, vascular endothelial growth factor B
					EST, Highly similar to 40S RIBOSOMAL PROTEIN S2 [M.musculus], EST, Highly similar to 40S RIBOSOMAL PROTEIN S2
					[R.norvegicus], EST, Moderately similar to S08228 ribosomal protein S2, cytosolic [H.sapiens], EST, Weakly
					similar to S08228 ribosomal protein S2, cytosolic [H.sapiens], ESTs, Highly similar to RS2_HUMAN 40S RIBOSOMAL PROTEIN S2
					[H.sapiens], ESTs, Moderately similar to 40S RIBOSOMAL PROTEIN S2 [M.musculus], repeat family 3 gene,
1959	10268	NM_031838	a		ribosomal protein S2

TABUE 8	REMINER 3	HOWOTOGNEVI	SMOTIATION		Aiiy. Dockei No. 44921-5089WO Doc. No. 1798897.1
809. ID No.	ldentifier	ConBank Acel Ref. Seq. ID No.	Model Gode	Homologous Gene, 1 Neme	Homologovs Glyster Name
					EST, Highly similar to 40S RIBOSOMAL PROTEIN S2 [M.musculus], EST, Highly similar to 40S RIBOSOMAL PROTEIN S2 [R.norvegicus], EST, Moderately similar to S08228 ribosomal protein S2, cytosolic [H.sapiens], EST, Weakly similar to S08228 ribosomal protein S2, cytosolic [H.sapiens], ESTs, Highly
					similar to RS2_HUMAN 40S RIBOSOMAL PROTEIN S2 [H.sapiens], ESTs, Moderately similar to 40S RIBOSOMAL PROTEIN S2 [M.musculus], repeat family 3 gene,
1959	10269	NM_031838	аа		ribosomal protein S2
					EST, Highly similar to 40S RIBOSOMAL PROTEIN S2 [M.musculus], EST, Highly similar to 40S RIBOSOMAL PROTEIN S2 [R.norvegicus], EST, Moderately similar to S08228 ribosomal protein S2, cytosolic [H.sapiens], EST, Weakly similar to S08228 ribosomal protein S2, cytosolic [H.sapiens], ESTs, Highly similar to RS2_HUMAN 40S RIBOSOMAL PROTEIN S2 [H.sapiens], ESTs, Moderately similar to 40S RIBOSOMAL PROTEIN S2 [M.musculus], repeat family 3 gene,
1959	10267	NM_031838	n,aa		ribosomal protein S2 expressed sequence AU022220, hypothetical protein FLJ21032, stearoyl-CoA desaturase (delta-9- desaturase), stearoyl-Coenzyme A desaturase 1, stearoyl-Coenzyme A desaturase 2, stearoyl-coenzyme A
1960	15077	NM_031841	b		desaturase 3
1961	16726	NM_031855	x	ketohexokinase, ketohexokinase (fructokinase)	ketohexokinase, ketohexokinase (fructokinase)
1962	25802	NM_031969	а	calmodulin 1, calmodulin 1 (phosphorylase kinase, delta)	
					Calmodulin III, ESTs, Highly similar to A Chain A, Calmodulin Complexed With Calmodulin-Binding Peptide From Smooth Muscle Myosin Light Chain Kinase (SUB 2-148 [H.sapiens], R.norvegicus CaMil retropseudogene (clone lambda SC27), RIKEN cDNA 2310068O22 gene, calmodulin, calmodulin 1, calmodulin 1 (phosphorylase kinase, delta),
1962	19191	NM 031969	С	calmodulin 1, calmodulin 1 (phosphorylase kinase, delta)	calmodulin 2, calmodulin 2 (phosphorylase kinase, delta), calmodulin 3, calmodulin-like 3

TABLES	e (HUMAN)	HOMOLOGUE AN	INOTATIONS	Thinks a state of the	Aily, Docket No. 44921-5039WO Doc. No. 1793897.1
839. ID No.	nelllimebl	ConBank Acel Ref. Seg. (D No.	Model Gode	Hemelogovs Genre Name	Honologous Cluster Name
				calmodulin 1, calmodulin 1 (phosphorylase kinase,	Calmodulin III, ESTs, Highly similar to A Chain A, Calmodulin Complexed With Calmodulin-Binding Peptide From Smooth Muscle Myosin Light Chain Kinase (SUB 2-148 [H.sapiens], R.norvegicus CaMII retropseudogene (clone lambda SC27), RIKEN cDNA 2310068O22 gene, calmodulin, calmodulin 1, calmodulin 1 (phosphorylase kinase, delta), calmodulin 2, calmodulin 2 (phosphorylase kinase, delta),
1962	19195	NM_031969	r	calmodulin 1, calmodulin 1 (phosphorylase kinase,	calmodulin 3, calmodulin-like 3 Calmodulin III, ESTs, Highly similar to A Chain A, Calmodulin Complexed With Calmodulin-Binding Peptide From Smooth Muscle Myosin Light Chain Kinase (SUB 2-148 [H.sapiens], R.norvegicus CaMII retropseudogene (clone lambda SC27), RIKEN cDNA 2310068O22 gene, calmodulin, calmodulin 1, calmodulin 1 (phosphorylase kinase, delta), calmodulin 2, calmodulin 2 (phosphorylase kinase, delta),
1962	19190	NM_031969	P	delta)	calmodulin 3, calmodulin-like 3 EST, Weakly similar to HHHU27 heat shock protein 27 [H.sapiens], ESTs, Moderately similar to HHHU27 heat shock protein 27 [H.sapiens].
1963	17734	NM_031970	v,General	heat shock 70kD protein 1B, heat shock protein, 70 kDa 1	crystallin, alpha C, heat shock 27kD protein 1, heat shock 27kD protein 3 ESTs, Weakly similar to BCHUIA S-100 protein alpha chain [H.sapiens], ESTs, Weakly similar to S10A MOUSE S-100 PROTEIN, ALPHA CHAIN [M.musculus], Homo sapiens cDNA FLJ10018 fis, clone HEMBA1000531, RIKEN cDNA B230217N24 gene, S100 calcium binding protein A1, S100 calcium binding protein A11, S100 calcium-binding protein A1, S100 calcium-binding protein A11 (calgizzarin), S100 calcium-binding protein A11 (calgizzarin), S100 calcium-binding protein P, expressed sequence A1266795, heat shock 70kD protein 1A, heat shock 70kD protein 1B
1965	15470	NM_031978	f	NDG I	proteasome (prosome, macropain) 26S subunit, non-ATPase, 1
1966	18502	NM_031984	c	calbindin 1, (28kD), calbindin-28K	ESTs, Moderately similar to CALBINDIN [M.musculus], calbindin 1, (28kD), calbindin-28K

Seg., (D) No. :		The Article of the State of the			Dos. No. 1793397.1
200102.002000	ldentifier	ConBank Ace./ Ref. Seq. (D No.)	Model Code	Honologous Cene	Homologous Gluster Name
1967	19768	NM_031986	v,aa,General		ESTs, Highly similar to APB1 RAT AMYLOID BETA A4 PRECURSOR PROTEIN-BINDING FAMILY A MEMBER 1 [R.norvegicus], Mus musculus, Similar to hypothetical protein, clone MGC:11704 IMAGE:3964815, mRNA, complete cds, RIKEN cDNA 2310008D10 gene, amyloid beta (A4) precursor protein-binding, family A, APBA1: amyloid beta (A4) precursor protein-binding, family A, member 1 (X11), amyloid beta (A4) precursor protein-binding, family A, member 1 (X11), amyloid beta (A4) precursor protein-binding, family A, member 1 (X11), amyloid beta (A4) precursor protein-binding, family A, member 3, syndecan binding protein, syndecan binding protein, syndecan binding protein (syntenin), syntenin-2 protein ESTs, Weakly similar to T42724 p190-B protein - mouse [M.musculus], PTPL1-associated RhoGAP 1, RIKEN
					cDNA 1700026N20 gene, RIKEN cDNA 1700026N20 gene, RIKEN cDNA 1700112L09 gene, chimerin (chimaerin) 2, minor histocompatibility antigen HA-1, oligophrenin 1, rho
1968	723	NM_032084	n		GTPase activating protein 5
1969	17935	NM 032615	а	membrane interacting protein of RGS16	hypothetical protein FLJ20207, membrane interacting protein of RGS16
1970	16831	NM_033095	n		
1971	25468	NM_033234	C,Z		
1971	25469	NM_033234	С		
1971	17832	NM_033234	с,р	hemoglobin beta chain complex, hemoglobin, beta hemoglobin beta chain	
1971	17829	NM 033234	c,z	complex, hemoglobin, beta	
				3000	ESTs, Highly similar to LDHH_HUMAN L-LACTATE DEHYDROGENASE H CHAIN [H.sapiens], Lactate dehydrogenease B, RIKEN cDNA 1700124B08 gene, lactate dehydrogenase 2, B chain, lactate dehydrogenase B, malate dehydrogenase 1, NAD (soluble),
1972	4723	NM_033235	Z		malate dehydrogenase, soluble
1973	1409	NM_033349	p,General	glyoxylase 2, hydroxyacyl glutathione hydrolase	Mus musculus, Similar to hydroxyacyl glutathione hydrolase, clone MGC:6697 IMAGE:3583919, mRNA, complete cds, RIKEN cDNA 1500017E18 gene, RIKEN cDNA 2810014I23 gene, RIKEN cDNA C330022E15 gene, hydroxyacyl glutathione hydrolase, hypothetical protein MGC2605, protein expressed in thyroid

TABLES	e HUMAN	HOMOTOGALE VA	SKOTATOK		Atty, Docket No. 44921-51199WO Doc. No. 1798997.1
	ldentifier	ConBenk Acol Ref. Seq. ID No.	Model Gode	Hamologous Cene Name	Homologous Glusia Name
1974	19998	NM 033352	General	PDZ domain containing 1	ESTs, Weakly similar to T30259 multiple PDZ domain protein - mouse [M.musculus], ESTs, Weakly similar to T46612 multi PDZ domain protein 1 - rat [R.norvegicus], PDZ domain containing 1, channel-interacting PDZ domain protein, hypothetical protein FLJ22756, multiple PDZ domain protein, semaF cytoplasmic domain associated protein 3, solute carrier family 9 (sodium/hydrogen exchanger), isoform 3 regulatory factor 1, solute carrier family 9 (sodium/hydrogen exchanger), isoform 3 regulatory factor 2, syntaxin binding protein 4
10.4		555552	Silvini	22 domain containing 1	ESTs, Moderately similar to S47073 finger protein HZF2, Krueppel-related [H.sapiens], ESTs, Moderately similar to T12489 hypothetical protein DKFZp572P0920.1 [H.sapiens], ESTs, Weakly similar to TC17 MOUSE TRANSCRIPTION FACTOR 17 [M.musculus], ESTs, Weakly similar to Z184_HUMAN ZINC FINGER PROTEIN 184 [H.sapiens], expressed sequence Al875089, transcription factor 17, transcription factor 17-like 1,
1975	1410	NM_052798	d	zinc finger protein 354A cysteine dioxygenase 1,	transcription factor 17-like 2 RIKEN cDNA 2900092E17 gene,
1976	15028	NM_052809	f	cytosolic, cysteine dioxygenase, type I	cysteine dioxygenase 1, cytosolic, cysteine dioxygenase, type I
1977	5176	NM_053297	и	pyruvate kinase 3, pyruvate kinase, muscle	
1075					EST, Moderately similar to S12583 polyubiquitin 4 - mouse [M.musculus], Homo sapiens UBBP2 pseudogene for ubiquitin UBB, RIKEN cDNA 2700054004 gene, diubiquitin, expressed sequence Al194771, expressed sequence AL033289,
1978	7660	NM_053299	i	Homer, neuronal immediate early gene, 3,	ubiquitin B, ubiquitin C CAT56 protein, EST, Weakly similar to A28996 proline-rich protein M14 precursor - mouse [M.musculus], EST, Weakly similar to JE0291 FB19 protein [H.sapiens], Homer, neuronal immediate early gene, 1B, RuvB-like protein 1, homer, neuronal immediate early gene, 1, homer, neuronal immediate early gene, 2, proline rich protein, proline rich protein 2, protein
1979	5117	NM_053310	р	early gene, 3	phosphatase 1, regulatory subunit 10 ESTs, Moderately similar to protein
1981	17473	NM 053319	a,v	dynein, cytoplasmic, light chain 1, dynein,	inhibitor of nitric oxide synthase [M.musculus], RIKEN cDNA 6720463E02 gene, Rattus norvegicus dynein light chain-2 (Dlc2) mRNA, complete cds, dynein, cytoplasmic, light chain 1, dynein, cytoplasmic, light polypeptide
				insulin-like growth factor binding protein, acid labile	Lawrence
1982	25480	NM_053329	g	subunit	

TABLE	r Humani	HOWOLOGUE W	SKONATOKI		Atty, Docket No. 44921-5139WO Doc, No. 1793397.1
899, ID No.	ldentifier	Cenbenk Acel Ref. Seq. ID No.	Model)Code	Homologous Cene Name	Homologous Cluster Name b
4082	24077	NIM 052220		insulin-like growth factor binding protein, acid labile	ESTs, Weakly similar to ALS RAT INSULIN-LIKE GROWTH FACTOR BINDING PROTEIN COMPLEX ACID LABILE CHAIN PRECURSOR [R.norvegicus], ESTs, Weakly similar to JC6128 insulin-like growth factor binding complex acid labile chain - mouse [M.musculus], ESTs, Weakly similar to membrane glycoprotein [M.musculus], KIAA0644 gene product, glycoprotein 1a, alpha polypeptide, hypothetical protein FLJ20156, insulin-like growth factor binding protein, acid labile subunit, nogo receptor, reticulon 4 receptor, toll-like receptor 6, tumor endothelial
1982	21977	NM_053329		subunit	marker 5 precursor EST, Moderately similar to 2113200B ribosomal protein L21 [H.sapiens], EST, Moderately similar to RL21 MOUSE 60S RIBOSOMAL PROTEIN L21 [M.musculus], EST, Weakly similar to 2113200B ribosomal protein L21 [H.sapiens], EST, Weakly similar to RL21 MOUSE 60S RIBOSOMAL PROTEIN L21 [M.musculus], EST, Weakly similar to RL21 [M.musculus], EST, Weakly similar to RL21 HUMAN 60S RIBOSOMAL PROTEIN L21 [H.sapiens], ESTs, Moderately similar to RL21 RAT 60S RIBOSOMAL PROTEIN L21 [R.norvegicus], RIKEN cDNA 2700085M18 gene, ribosomal
1983	14929	NM_053330	e,General	ribosomal protein L21	protein L21 EST, Moderately similar to 2113200B ribosomal protein L21 [H.sapiens], EST, Moderately similar to RL21 MOUSE 60S RIBOSOMAL PROTEIN L21 [M.musculus], EST, Weakly similar to 2113200B ribosomal protein L21 [H.sapiens], EST, Weakly similar to RL21 MOUSE 60S RIBOSOMAL PROTEIN L21 [M.musculus], EST, Weakly similar to RL21 HUMAN 60S RIBOSOMAL PROTEIN L21 [H.sapiens], ESTs, Moderately similar to RL21 RAT 60S RIBOSOMAL PROTEIN L21 [H.sapiens], ESTs, Moderately similar to RL21 RAT 60S RIBOSOMAL PROTEIN L21 [R.norvegicus], RIKEN cDNA 2700085M18 gene, ribosomal protein L21
1984	16407	NM_053332	c,e	cubilin (intrinsic factor- cobalamin receptor)	DNA segment, Chr 2, Wayne State University 88, expressed, EST, Weakly similar to T09456 intrinsic factor-B12 receptor Cubilin precursor [H.sapiens], ESTs, Moderately similar to T09456 intrinsic factor-B12 receptor Cubilin precursor [H.sapiens], Homo sapiens cDNA FLJ12558 fis, clone NT2RM4000787, bone morphogenetic protein 1, cubilin (intrinsic factor- cobalamin receptor), expressed sequence AL022750, platelet-derived growth factor, C polypeptide, tolloid- like, tolloid-like 2, tumor necrosis factor induced protein 6, tumor necrosis factor, alpha-induced protein 6

TABLES	RAMUUH R	HOMOLOGUEAN	INOTATIONS		XXXY: Docket No. 44921-5039WO Doc. No. 1793397.1
Seq. [D] No. : :	lideniliter	GenBank Ace./ Ref. Seq. ID No.	Model Code	Homologovs Gene Mane	Homologous Cluster Name
1985	15790	NM 053341	j,×	chromosome 19 open reading frame 3, regulator of G-protein signaling 19 interacting protein 1	chromosome 19 open reading frame 3, hypothetical protein FLJ20075
					ESTs, Weakly similar to CGHU2S collagen alpha 2(I) chain precursor [H.sapiens], KIAA1026 protein, RIKEN cDNA 1110030G05 gene, RIKEN cDNA 9030409G11 gene, collagen, type I, alpha 1, collagen, type I, alpha 2, collagen, type II, alpha 1 (primary osteoarthritis, spondyloepiphyseal dysplasia, congenital), hypothetical protein FLJ20654, nischarin, procollagen, type I, alpha 1,
1986	6154	NM 053356	p	collagen, type I, alpha 2, procollagen, type I, alpha 2	procollagen, type I, alpha 2, procollagen, type II, alpha 1
1987	9215	NM_053374	i	proconagen, type i, aipna z	interleukin 18 binding protein ESTs, Weakly similar to NPT2 RAT RENAL SODIUM-DEPENDENT PHOSPHATE TRANSPORT PROTEIN 2 [R.norvegicus], ESTs, Weakly similar to NPT2_HUMAN RENAL SODIUM- DEPENDENT PHOSPHATE TRANSPORT PROTEIN 2 [H.sapiens], Homo sapiens, Similar to solute carrier family 34 (sodium phosphate), member 1, clone MGC:18179 IMAGE:4155326, mRNA, complete cds, Rattus norvegicus mRNA for NaPi-2 alpha, complete cds, Solute carrier family 17 (sodium/hydrogen exchanger), member 2, expressed sequence Al649385, solute carrier family 34 (sodium phosphate), member 1, solute carrier family 34 (sodium phosphate),
1988	6416	NM_053380	General	small muscle protein, X-	member 2
1989	19113	NM_053395	a		small muscle protein, X-linked ESTs, Highly similar to FMO3_HUMAN DIMETHYLANILINE MONOOXYGENASE [H.sapiens], Flavin-containing monooxygenase 1, Homo sapiens DNA sequence from PAC 127D3 on chromosome 1q23-25. Contains FMO2 and FMO3 genes for Flavin-containing Monooxygenase 2 and Flavin-containing Monooxygenase 3 (Dimethylaniline Monooxygenase (N-Oxide 3, EC1.14.13.8, Dimethylaniline Oxidase 3, FMO II, FMO 3), and a gene for another, unknown, Flavin- containing Monooxygenase family protein. Contains ESTs and GSSs, flavin containing monooxygenase 1, flavin containing monooxygenase 2, flavin containing monooxygenase 3,
1990	2242	NM_053433	n,General		hypothetical protein PRO1257 RIKEN cDNA 1700065B19 gene,
1991	5561	NM_053438	у		RIKEN cDNA 5730408C10 gene, zinc finger protein 103, zinc finger protein homologous to Zfp103 in mouse

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		HOMOFOGATEV	SKOTEATOKE		Aity. Doctot No. 44921-5039W6 Doc. No. 1793397.1
809. ID No.		Cenbenk Ace./ Ref. Seq. ID No.	Model Code	Homologovs Cenio Name	Homologous Gluster Name
1992	14670	NM 053439	n,General		ESTs, Highly similar to RAB7 RAT RAS-RELATED PROTEIN RAB-7 [R.norvegicus], RAB7, member RAS oncogene family, RAN, member RAS oncogene family, RIKEN cDNA 1700009N14 gene
				stathmin-like 2, superiorcervical ganglia,	stathmin-like 2, stathmin-like 4, superiorcervical ganglia, neural
1993	17102	NM_053440	w	neural specific 10	specific 10 ESTs, Weakly similar to 1615347A ras p21 GTPase activating protein [M.musculus], KIAA1938 protein, Mus musculus, Similar to RAS p21 protein activator, clone MGC:7759 IMAGE:3498774, mRNA, complete cds, RAS protein activator like 2, Rattus norvegicus DOC2/DAB2 interactive protein mRNA, complete
1994	24762	NM_053442	General		cds, expressed sequence BB079060, hypothetical protein FLJ21438
					ESTs, Moderately similar to RGS8 RAT REGULATOR OF G-PROTEIN SIGNALING 8 [R.norvegicus], regulator of G-protein signaling 18, regulator of G-protein signaling 2, regulator of G-protein signaling 13, regulator of G-protein signalling 13, regulator of G-protein signalling 2,
1995	8085	NM_053453	General		24kD, regulator of G-protein signalling 8 NEFA precursor, expressed sequence
1996	4622	NM_053463	d		Al607786, nucleobindin, nucleobindin 1, nucleobindin 2
1997	21866	NM 053472	p	cytochrome c oxidase subunit IV isoform 2, cytochrome c oxidase, subunit IVb	EST, Weakly similar to COX4_HUMAN CYTOCHROME C OXIDASE POLYPEPTIDE IV PRECURSO [H.sapiens], cytochrome c oxidase subunit IV, cytochrome c oxidase subunit IV isoform 2, cytochrome c oxidase, subunit IV, cytochrome c oxidase, subunit IVa, cytochrome c oxidase, subunit IVb, expressed sequence AL024441
1998	9573	NM 053475	lh.	GOOD TO THE TOTAL THE TOTAL TO THE TOTAL TO THE TOTAL TO THE TOTAL TO THE TOTAL TO THE TOTAL TO THE TOTAL TO THE TOTAL TO THE TOTAL TO THE TOTAL TOT	protein tyrosine phosphatase 4a1, protein tyrosine phosphatase 4a2, protein tyrosine phosphatase type IVA, member 1, protein tyrosine phosphatase type IVA, member 2, protein tyrosine phosphatase type IVA, member 3
					DNA polymerase alpha 2, 68 kDa, Mus musculus, Similar to DNA polymerase alpha 2, 68 kDa, clone MGC:11533 IMAGE:3602559, mRNA, complete cds, expressed sequence Al573378, polymerase (DNA-directed), alpha
1999	16137	NM_053480	k		(70kD) ESTs, Weakly similar to A Chain A, Importin Alpha, Mouse [M.musculus], expressed sequence AW146299, karyopherin (importin) alpha 2, karyopherin alpha 2 (RAG cohort 1,
2000	15556	NM_053483	у		importin alpha 1) EST, Moderately similar to CALCYCLIN [R.norvegicus], S100
2001	16394	NM_053485	General		calcium-binding protein A6 (calcyclin), calcium binding protein A6 (calcyclin)

		HONOFOGUE AV	INOTATIONS		Aity. Docto: No. 44921-5039WO Doc. No. 1793897.1
Seq. [D	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	ConBank Ace./ Ref. Sog. ID No.	Model Gode '-	Honologous Gene Neme	Homologous Cluster Name 4
2002	4290	NM_053487	j.y		peroxisomal biogenesis factor 11A, peroxisomal biogenesis factor 11B
					EST, Moderately similar to Y025_HUMAN HYPOTHETICAL PROTEIN KIAA0025 [H.sapiens], RIKEN cDNA 5031400M07 gene, homocysteine-inducible, endoplasmic reticulum stress-inducible, ubiquitin-like domain member 1, hypothetical
2004	18826	NM_053523	d		protein FLJ22313 DEAD/H (Asp-Glu-Ala-Asp/His) box
					polypeptide, Y chromosome, DNA segment, Chr 1, Pasteur Institute 1, ESTs, Moderately similar to DDXY_HUMAN DEAD BOX PROTEIN 3, Y-CHROMOSOMAL [H.sapiens], KIAA0801 gene product, RNA helicase, Rattus norvegicus RNA helicase with arginine-serine-rich domain mRNA, complete cds, expressed sequence Al324246, expressed sequence Al325430,
2005	7764	NM_053525	aa		expressed sequence C86129
2006	14199	NM 053538	c		Lysosomal-associated multispanning membrane protein-5, lysosomal-associated protein transmembrane 5
2007	1058	NM_053539	c,d		
					DNA segment, Chr 17, human D6S81E 1, EST, Weakly similar to HE47 RAT PROBABLE ATP- DEPENDENT RNA HELICASE P47 [R.norvegicus], HLA-B associated transcript 1, KIAA0111 gene product, Mus musculus, clone MGC:6664 IMAGE:3498954, mRNA, complete cds, RIKEN cDNA 2610307C23 gene, eukaryotic translation initiation factor 4A, isoform 2, eukaryotic translation initiation factor 4A1, eukaryotic translation initiation factor 4A2, nuclear RNA helicase, DECD variant of DEAD
2008	4327	NM_053563	General		box family
					ESTs, Weakly similar to JE0096 myocilin - mouse [M.musculus], Homo sapiens NOE3-4 (NOE3) mRNA, complete cds, alternatively spliced, expressed sequence AW742568, olfactomedin related ER localized
2009	1342	NM_053573	h		protein ESTs, Moderately similar to
2010	19254	NM_053576	h,s	selenium glutathione	AOP2_HUMAN ANTIOXIDANT PROTEIN 2 [H.sapiens], anti-oxidant protein 2 (non-selenium glutathione peroxidase, acidic calcium- independent phospholipase A2), peroxiredoxin 5
2010	19253	NM_053576	h	selenium glutathione	ESTs, Moderately similar to AOP2_HUMAN ANTIOXIDANT PROTEIN 2 [H.sapiens], anti-oxidant protein 2 (non-selenium glutathione peroxidase, acidic calcium- independent phospholipase A2), peroxiredoxin 5

TABLE	88 HUMAN	HOMOLOGUEA	NNOTATIONS		Atty. Docket No. 44921-5039W0 Doc. No. 1793897.1
Seg. (D No.	ldentifier	Condonk Ace/ Ref. Seq. ID No.	Model(Gode)	Homologous Cens	Homologous Cluster Name
2011	3049	NM_053582	p,cc,General		ESTs, Highly similar to JC7189 tubulointerstitial nephritis antigen [H.sapiens], P3ECSL, cathepsin B, lipocalin 7, tubulointerstitial nephritis antigen
					ESTs, Highly similar to JC7189 tubulointerstitial nephritis antigen [H.sapiens], P3ECSL, cathepsin B, lipocalin 7, tubulointerstitial nephritis
2011	21423	NM_053582	o,General		antigen EST, Moderately similar to CYTOCHROME C OXIDASE POLYPEPTIDE VB PRECURSOR [R.norvegicus], cytochrome c oxidase subunit Vb, cytochrome c oxidase, subunit Vb
2013	21445	NM 053587	t,v	 	Suburit VD
2014	20871	NM 053591			ESTs, Weakly similar to MDP1 MOUSE MICROSOMAL DIPEPTIDASE PRECURSOR [M.musculus], ESTs, Weakly similar to S33757 membrane dipeptidase [M.musculus], RIKEN cDNA 1700018F16 gene, dipeptidase, putative
			<i>y.</i>		metallopeptidase (family M19) ESTs, Weakly similar to MDP1 MOUSE MICROSOMAL DIPEPTIDASE PRECURSOR [M.musculus], ESTs, Weakly similar to S33757 membrane dipeptidase [M.musculus], RIKEN cDNA 1700018F16 gene, dipeptidase 1 (renal), putative dipeptidase, putative
2014	20870	NM_053591	1		metallopeptidase (family M19) protein tyrosine phosphatase, receptor
2015	21044	NM_053594 NM_053596	d		type, R KIAA0604 gene product, endothelin converting enzyme 1, endothelin converting enzyme-like 1, expressed sequence AW322500, expressed sequence BB127715, mel transforming oncogene-like 1, membrane metallo endopeptidase
2016	21708	NM_053596	z		KIAA0604 gene product, endothelin converting enzyme 1, endothelin converting enzyme-like 1, expressed sequence AW322500, expressed sequence BB127715, mel transforming oncogene-like 1, membrane metallo endopeptidase
				nuclear proten 1, p8 protein (candidate of	ESTs, Weakly similar to Gene product
2017	1597	NM_053611	t	metastasis 1)	with similarity to Rat P8 [H.sapiens]
				Bardet-Biedl syndrome 2, Bardet-Biedl syndrome 2	
2018	5565	NM_053618	General	(human) fatty acid-Coenzyme A ligase, long chain 4, fatty- acid-Coenzyme A ligase,	
2019	13004	NM_053623	t	long-chain 4	1

TABLES	S HUMAN	HOMOLO@UE AN	SKIOLITATION		Atty: Docket No. 44921-5039WO Doc. No. 1793897.1
Seq. ID No.	ldentliter	ConDenk Ace./ Ref. Soq. ID No.	Model Gode	Homologous Come Name	Homologous Austa Name
					D-amino acid oxidase, D-amino-acid oxidase, D-aspartate oxidase, EST, Weakly similar to OXDA RAT D-AMINO ACID OXIDASE [R.norvegicus], ESTs, Highly similar to OXDA RAT D-AMINO ACID OXIDASE [R.norvegicus], RIKEN cDNA 5330420D20 gene, RIKEN cDNA
2020	1127	NM_053626	g	beta-carotene 15, 15'-	5730402C02 gene EST, Moderately similar to 0806162D
2021	18644	NM 053648	ln	dioxygenase, beta- carotene 15,15'- dioxygenase	protein COII [M.musculus], EST, Weakly similar to 810024D cytochrome oxidase II [H.sapiens]
				a.o.ygo.iaoo	c-fos induced growth factor, c-fos induced growth factor (vascular endothelial growth factor D), vascular endothelial growth factor, vascular endothelial growth factor B, vascular
2022	21637	NM_053653	<u>р</u>		endothelial growth factor C ESTs, Highly similar to CG1C RAT
2023	3454	NM 053662	cc		G1/S-SPECIFIC CYCLIN C [R.norvegicus], Homo sapiens, clone IMAGE:3537447, mRNA, partial cds, RIKEN cDNA 1810009O10 gene, cyclin C, cyclin K, cyclin L, cyclin L ania-6a, cyclin T2
					with Glu/Asp-rich carboxy-terminal domain, 2, Cbp/p300-interacting transactivator, with Glu/Asp-rich carboxy-terminal domain, 4, ESTs, Weakly similar to MRG1_HUMAN MSG-RELATED PROTEIN 1 [H.sapiens], expressed sequence
2024	16121	NM_053698	h.j.z		AW742964 with Glu/Asp-rich carboxy-terminal domain, 2, Cbp/p300-interacting transactivator, with Glu/Asp-rich carboxy-terminal domain, 4, ESTs, Weakly similar to MRG1_HUMAN MSG-RELATED PROTEIN 1 [H.sapiens], expressed sequence
2024	16122 25379	NM_053698 NM_053713	h,j,z General		AW742964
0005	40000	LUA 050740			ESTs, Moderately similar to CPBP RAT CORE PROMOTER ELEMENT- BINDING PROTEIN [R.norvegicus], Kruppel-like factor 4 (gut), RIKEN cDNA 7420700M05 gene, core
2025 2026	13622 15376	NM_053713 NM_053747	General h	ubiquilin 1	promoter element binding protein
2027	1218	NM_053748	b		expressed sequence C86324, hypothetical protein FLJ23590 cytochrome P450, 40 (25- hydroxyvitamin D3 1 alpha-
2028	1137	NM_053763	у		hydroxylase), cytochrome P450, subfamily XXVIIB (25-hydroxyvitamin D-1-alpha-hydroxylase), polypeptide 1 MKP-1 like protein tyrosine
2029	15996	NM_053769	сс	dual specificity phosphatase 1, protein tyrosine phosphatase, non- receptor type 16	phosphatase, dual specificity phosphatase 1, dual specificity phosphatase 13, dual specificity phosphatase 14, dual specificity phosphatase 2, expressed sequence

TABLE 8	R HIMIMAN	HOMOLOGUE AN	ROTATIONS		Ally, Docket No. 44921-5039W0 Doc. No. 1793397.1
Seg, Îd No.	ldenlifter	Cenbenk Acel Ref. Seq. ID No.	Model Gode in	Homologous Cane Name 1. 2	Homologous Clusier Name
2030	8652	NM_053774	g	ubiquitin specific protease	KIAA1453 protein, RIKEN cDNA 4930511O11 gene, expressed sequence AA409661, ubiquitin specific protease 2, ubiquitin specific protease 8
2031	14664	NM_053806	General		
2032	4361	NM_053812	k		B cell lymphoma 2 like, BCL2- antagonist/killer 1, BCL2-like 1, Bcl-w protein, Bcl2-like, Mus musculus N- BAK1 (Bak1) mRNA, complete cds, alternatively spliced, RIKEN cDNA 0610031G08 gene
					EST, Moderately similar to TIM1 RAT METALLOPROTEINASE INHIBITOR 1 PRECURSOR [R.norvegicus], EST, Weakly similar to TIM1 RAT METALLOPROTEINASE INHIBITOR 1 PRECURSOR [R.norvegicus], tissue inhibitor of metalloproteinase, tissue inhibitor of metalloproteinase 1 (erythroid potentiating activity,
2034	15002	NM 053819	b,x,bb,General		collagenase inhibitor)
					EST, Moderately similar to TIM1 RAT METALLOPROTEINASE INHIBITOR 1 PRECURSOR [R.norvegicus], EST, Weakly similar to TIM1 RAT METALLOPROTEINASE INHIBITOR 1 PRECURSOR [R.norvegicus], tissue inhibitor of metalloproteinase, tissue inhibitor of metalloproteinase 1 (erythroid potentiating activity,
2034	15003	NM_053819	b,I,x,bb,General		collagenase inhibitor)
2035	16173	NM_053822	t		
2036	17154	NM_053835	j,z		
2037	20868	NM_053843	t		
2037	20869	NM_053843	t	"	
2040	714	NM_053863	у		ESTs, Highly similar to CNT1_HUMAN SODIUM/NUCLEOSIDE COTRANSPORTER 1 [H.sapiens], ESTs, Moderately similar to A54892 Na+-dependent nucleoside transport protein cNT1 - rat [R.norvegicus]
					ESTs, Moderately similar to DUS6 RAT DUAL SPECIFICITY PROTEIN PHOSPHATASE 6 [R.norvegicus], ESTs, Weakly similar to DUS6 RAT DUAL SPECIFICITY PROTEIN PHOSPHATASE 6 [R.norvegicus], dual specificity phosphatase 10, dual specificity phosphatase 13, dual specificity phosphatase 14, dual specificity phosphatase 6, dual specificity phosphatase 6, dual specificity phosphatase 9, expressed sequence BB104621, mitogen- activated protein kinase phosphatase x, protein tyrosine phosphatase, non-
2041	19781	NM 053883	b		receptor type 16

TABLES	B: CHUMPAN	HOMOFOGAE VA	- exolitations		Atty. Docket No. 44921-5039WO Doc. No. 1793397.1
Seq. ID No.	ldentiller	Conbank Ace./ Ref. Seq. (D No.		Homologous Cene Name	Homologous Cluster Name
2041	19780	NM_053883	Ь		ESTs, Moderately similar to DUS6 RAT DUAL SPECIFICITY PROTEIN PHOSPHATASE 6 [R.norvegicus], ESTs, Weakly similar to DUS6 RAT DUAL SPECIFICITY PROTEIN PHOSPHATASE 6 [R.norvegicus], dual specificity phosphatase 10, dual specificity phosphatase 13, dual specificity phosphatase 14, dual specificity phosphatase 6, dual specificity phosphatase 9, expressed sequence BB104621, mitogenactivated protein kinase phosphatase x, protein tyrosine phosphatase, non-receptor type 16
2042	1454	NM_053887	General		ESTs, Moderately similar to CD5R
2043	1660	NM_053891	g		MOUSE CYCLIN-DEPENDENT KINASE 5 ACTIVATOR 1 PRECURSOR [R.norvegicus], cyclin- dependent kinase 5, regulatory subunit (p35), cyclin-dependent kinase 5, regulatory subunit 1 (p35), cyclin- dependent kinase 5, regulatory subunit 2 (p39)
2044	712	NM_053896	k		coagulation factor II (thrombin)
2045	753	NM_053897	k		receptor-like 1, coagulation factor II (thrombin) receptor-like 2
2046	794	NM_053902	General		RIKEN cDNA 4432411A05 gene, kynureninase (L-kynurenine hydrolase) ESTs, Weakly similar to
2047	17937	NM_053911	f		ARNO_HUMAN ARF NUCLEOTIDE- BINDING SITE OPENER [H.sapiens]
					DNA segment, Chr 10, ERATO Doi 398, expressed, ESTs, Weakly similar to PTNL RAT PROTEIN TYROSINE PHOSPHATASE, NON-RECEPTOR TYPE 21 [R.norvegicus], Homo sapiens, Similar to erythrocyte membrane protein band 4.1-like 3, clone MGC:12343 IMAGE:4044866, mRNA, complete cds, Mus musculus adult male pituitary gland cDNA, RIKEN full-length enriched library, clone:5330430110, full insert sequence, Rattus norvegicus protein tyrosine phosphatase 2E (PTP2E) mRNA, complete cds, erythrocyte membrane protein band 4.1 (elliptocytosis 1, RH-linked), erythrocyte membrane protein band 4.1-like 1, erythrocyte protein band 4.1-like 3, erythrocyte protein band 4.1-like 3, erythrocyte
2048	8188	NM_053927	General		protein band 4.1-like 3, protein tyrosine phosphatase, non-receptor type 21

TABLES	KAMUH)	HOMOFOGÁE W	i enoltatoni	· · · · · · · · · · · · · · · · · · ·	Alily, Docker No. 44921-51189WO Doc. No. 1793397.1
809. [D No.	Mendi	Cenbenk Ace./ Rei. Seg. ID No.	Model Gode 4	Homologous Gene Name	
					endothelial differentiation, lysophosphatidic acid G-protein- coupled receptor 4, endothelial differentiation, lysophosphatidic acid G- protein-coupled receptor 7, endothelial differentiation, lysophosphatidic acid G- protein-coupled receptor, 2, endothelial differentiation, lysophosphatidic acid G-protein- coupled receptor, 4, endothelial differentiation, lysophosphatidic acid G- protein-coupled receptor, 7, putative G
2050	1628 13954	NM_053936 NM_053955	h General		protein-coupled receptor snGPCR32 ESTs, Highly similar to B46290 mu-
2052	408	NM 053961	General		crystallin [H.sapiens], crystallin, mu
2032	400	14W_033901	General		DNA segment, Chr 9, ERATO Doi 85, expressed, Homo sapiens aconitase precursor (ACON) mRNA, nuclear gene encoding mitochondrial protein, partial cds, RIKEN cDNA 5031409G22 gene, aconitase 1, aconitase 1, soluble, aconitase 2, mitochondrial, iron-responsive element-binding
2052	19991	NM 053961	a	/	protein
2052	16190	NM_053961	q		carcinoma-associated antigen 64 (HCA64) mRNA, complete cds, RIKEN cDNA 1300014E15 gene, RIKEN cDNA 1300017C12 gene, RIKEN cDNA 1810022C23 gene, RIKEN cDNA 2610009M20 gene, RIKEN cDNA 4933417A18 gene, enoyl Coenzyme A hydratase, short chain, 1, mitochondrial, peroxisomal D3,D2- enoyl-CoA isomerase, peroxisomal delta3, delta2-enoyl-Coenzyme A isomerase
2052	21355	NM_053961	j,l,y,z		
2055	15136 15135	NM_053971 NM_053971	aa		EST, Moderately similar to I51803 TAXREB107 [H.sapiens], ESTs, Highly similar to I51803 TAXREB107 [H.sapiens], ribosomal protein L6 EST, Moderately similar to I51803 TAXREB107 [H.sapiens], ESTs, Highly similar to I51803 TAXREB107 [H.sapiens], ribosomal protein L6
2056	1764	NM_053974	h		translation initiation factor elF-4E [H.sapiens], RIKEN cDNA 2700069E09 gene, eukaryotic translation initiation factor 4E ADP-ribosylation factor related protein
2057	1292	NM 053980			1, RIKEN cDNA 1500006101 gene
					EST, Highly similar to 40S RIBOSOMAL PROTEIN S15A [R.norvegicus], EST, Weakly similar to RS1A_HUMAN 40S RIBOSOMAL PROTEIN S15A [R.norvegicus], ESTs, Weakly similar to RS1A_HUMAN 40S RIBOSOMAL PROTEIN S15A
2058	15468	NM_053982 NM_053985	q General		[H.sapiens] ESTs, Highly similar to HISTONE H3.3 [R.norvegicus], H3 histone, family 3A, H3 histone, family 3B, H3 histone, family 3B (H3.3B)

TABLE		HOMOLOGUE AV	SMOTATIONS		Ally. Docket No. 44921-5039W0 Doc. No. 1793397.1
(10 Seg. (D	ldenilliter	Cenbenk Ace./ Ref. Seg. ID No.	Model Code	Homologous Comp Namo:	Homologous Cluster Name
					CD36 antigen (collagen type I receptor, thrombospondin receptor)-like 2 (lysosomal integral membrane protein II), EST, Moderately similar to
2060	21066	NM_054001	t		LYII_HUMAN LYSOSOME MEMBRANE PROTEIN II [H.sapiens]
2061	17326	NM_054008	o		RGC32 protein, RIKEN cDNA 1190002H23 gene
2061	17327	NM_054008	cc		RGC32 protein, RIKEN cDNA 1190002H23 gene
2061	17329	NM_054008	g,o,cc		RGC32 protein, RIKEN cDNA 1190002H23 gene
2062	25253	NM 057099	j,l,m,p,z	proteasome (prosome, macropain) subunit, beta type 6, proteasome (prosome, macropain) subunit, beta type, 6	EST, Weakly similar to S17522 multicatalytic endopeptidase complex [H.sapiens], ESTs, Weakly similar to PROTEASOME DELTA CHAIN PRECURSOR [R.norvegicus], proteasome (prosome, macropain) subunit, beta type 6, proteasome (prosome, macropain) subunit, beta type 7, proteasome (prosome, macropain) subunit, beta type, 6, proteasome (prosome, macropain) subunit, beta type, 6, proteasome (prosome, macropain) subunit, beta type, 7
				proteasome (prosome, macropain) subunit, beta type 6, proteasome (prosome, macropain)	EST, Weakly similar to S17522 multicatalytic endopeptidase complex [H.sapiens], ESTs, Weakly similar to PROTEASOME DELTA CHAIN PRECURSOR [R.norvegicus], proteasome (prosome, macropain) subunit, beta type 6, proteasome (prosome, macropain) subunit, beta type 7, proteasome (prosome, macropain) subunit, beta type, 6, proteasome (prosome, macropain)
2062	22849	NM_057099	j,l	subunit, beta type, 6 A kinase (PRKA) anchor	subunit, beta type, 7
2063	19657	NM_057103	b,cc	protein (gravin) 12 UDP glycosyltransferase 1 family, polypeptide A cluster, UDP glycosyltransferase 1 family, polypeptide A6, UDP- glucuronosyltransferase 1	
2064	5492	NM_057105	w	family, member 1 UDP glycosyltransferase 1 family, polypeptide A cluster, UDP glycosyltransferase 1 family, polypeptide A6, UDP- glucuronosyltransferase 1	
2064	15126	NM_057105	r	family, member 1 UDP glycosyltransferase 1 family, polypeptide A cluster, UDP- glucuronosyltransferase 1	
2064	15125 15391	NM_057105 NM_057114	s n	family, member 1	EST, Moderately similar to TDX2_HUMAN THIOREDOXIN PEROXIDASE 2 [H.sapiens], EST, Weakly similar to TDX2_HUMAN THIOREDOXIN PEROXIDASE 2 [H.sapiens], peroxiredoxin 1

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Seq. ID No.	ldentifier	ConBonk Aced Ref. Seq. ID No.	Model Gode	Homologous Cene Name	Homologous Auster Name
2067	727	NM_057123 NM_057124	m		BCS1 (yeast homolog)-like, ESTs, Highly similar to PRS4_HUMAN 26S PROTEASE REGULATORY SUBUNIT 4 [H.sapiens], ESTs, Weakly similar to A44468 26S proteasome regulatory chain 4 [H.sapiens], expressed sequence Al325227, protease (prosome, macropain) 26S subunit, ATPase 1, proteasome (prosome, macropain) 26S subunit, ATPase, 1 ESTs, Weakly similar to P2UR MOUSE P2U PURINOCEPTOR 1 [M.musculus], G protein-coupled receptor 35, purinergic receptor P2Y, G-protein coupled, 4, pyrimidinergic receptor P2Y, G-protein coupled, 4, pyrimidinergic receptor P2Y, G-protein coupled, 6
2069	15151	NM_057131	k		
2070	1892	NM_057144	ь		cysteine and glycine-rich protein 3 (cardiac LIM protein), cysteine rich protein, cysteine-rich protein 2, cysteine-rich protein 3, thymus LIM protein
2071	12333	NM_057155	f		
2071 2071	12331 12332	NM_057155 NM_057155	v,General f,General		
2072	17477	NM_057194	a,General		rich protein PRB3S [H.sapiens], EST, Weakly similar to CGHU3B collagen alpha 3(IV) chain precursor, long splice form [H.sapiens], EST, Weakly similar to D40750 proline-rich protein PRB1/2S [H.sapiens], EST, Weakly similar to JE0284 Mm-1 cell derived transplantability-associated protein 1b [H.sapiens], galectin-related inter-fiber protein, murine leukemia viral (bmi-1) oncogene homolog, phospholipid scramblase 1, phospholipid scramblase 2, phospholipid scramblase 3 2,4-dienoyl CoA reductase 1,
2073	15408	NM_057197	p,t		mitochondrial, 2,4-dienoyl CoA reductase 2, peroxisomal, peroxisomal trans 2-enoyl CoA reductase; putative short chain alcohol dehydrogenase
2073	15409	NM_057197	t		2,4-dienoyl CoA reductase 1, mitochondrial, 2,4-dienoyl CoA reductase 2, peroxisomal, peroxisomal trans 2-enoyl CoA reductase; putative short chain alcohol dehydrogenase
2074	7866	NM_057198	h		ESTs, Highly similar to PUR1_HUMAN AMIDOPHOSPHORIBOSYLTRANSFE RASE PRECURSOR [H.sapiens], ESTs, Moderately similar to PUR1_HUMAN AMIDOPHOSPHORIBOSYLTRANSFE RASE PRECURSOR [H.sapiens], RIKEN cDNA 5730454C12 gene, expressed sequence AA675351, expressed sequence C79945, glutamine fructose-6-phosphate transaminase 2, glutamine-fructose-6-phosphoribosyl pyrophosphate amidotransferase

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	neillineel	CenBank Ace/ Ref. Seq. ID No.	Model Gode	Homologous Gene Name	
2075 2076	14125 1743	NM_057208 NM_057210	h.j,y,z k,s	-	ESTs, Highly similar to A25530 tropomyosin, fibroblast [H.sapiens]
2077	10498	NM_078617	а		EST, Moderately similar to RS23_HUMAN 40S RIBOSOMAL PROTEIN S2 [R.norvegicus], EST, Weakly similar to RS23_HUMAN 40S RIBOSOMAL PROTEIN S2 [R.norvegicus], ESTs, Weakly similar to RS23_HUMAN 40S RIBOSOMAL PROTEIN S2 [R.norvegicus], Mus musculus, Similar to mitochondrial ribosomal protein S12, clone MGC:13892 IMAGE:4209358, mRNA, complete cds, RIKEN cDNA 2410044J15 gene, expressed sequence Al327385, mitochondrial ribosomal protein S12, ribosomal protein S23
2078	8820	NM_080399	n		
2079 2079	15701 20105	NM_080581 NM_080581	j,m,y,z		ATP-binding cassette, sub-family C (CFTR/MRP), member 1 (multiple drug resistance-associated protein), ATP-binding cassette, sub-family C (CFTR/MRP), member 1a, ATP-binding cassette, sub-family C (CFTR/MRP), member 1b, ATP-binding cassette, sub-family C (CFTR/MRP), member 2, ATP-binding cassette, sub-family C (CFTR/MRP), member 3, ATP-binding cassette, sub-family C (CFTR/MRP), member 3, ATP-binding cassette, sub-family C (CFTR/MRP), member 6, ESTs, Moderately similar to JE0336 canalicular multispecific organic anion transporter [H.sapiens], RIKEN cDNA 1700019L09 gene
2080	16109	NM 080585	c		
2081	1757	NM 080766	d		
2082	7108	NM_080778	у		ESTs, Highly similar to COT2 RAT COUP TRANSCRIPTION FACTOR 2 [R.norvegicus], Homo sapiens cDNA: FLJ22189 fis, clone HRC01043, RIKEN cDNA 2700033K02 gene, nuclear receptor subfamily 2, group E, member 3, nuclear receptor subfamily 2, group F, member 2
2083	132	NM_080782	k		cyclin-dependent kinase inhibitor 1A (P21), cyclin-dependent kinase inhibitor 1A (p21, Cip1), cyclin-dependent kinase inhibitor 1B (p27, kip1), cyclin-dependent kinase inhibitor 1B (p27, Kip1) cyclin-dependent kinase inhibitor 1A (P21), cyclin-dependent kinase
2083	133	NM_080782	1		inhibitor 1A (p21, Cip1), cyclindependent kinase inhibitor 1B (p27, kip1), cyclindependent kinase inhibitor 1B (p27, Kip1), cyclindependent kinase inhibitor 1B (p27, Kip1) HT014, Homo sapiens thioredoxindelta 3 (TXN delta 3) mRNA, partial cds, RIKEN cDNA 4930429J24 gene, expressed sequence AU021712,
2084	20122	NM_080887	General		thioredoxin, thioredoxin domain- containing 2 (spermatozoa), thioredoxin-like (32kD), thioredoxin- like, 32kD

TABLE E	KAMMUH X	HOMOLOGUE AV	SKOTKATOKE		Atty. Docket No. 44921-5139WO Doc. No. 1793897.1
Seg. 1D No.	lientifier	ConBenk Ace./ Rgf. Seq. ID No.	4 abs9 lebem	Homologous Cene Name	Homologous Gluster Name
2085 2086	6143 19952	NM_080892 NM_080902	e h		Homo sapiens, Similar to selenium binding protein 1, clone MGC:17268 IMAGE:4155238, mRNA, complete cds, selenium binding protein 1, selenium binding protein 2
2087	17546	NM 130401	b		epithelial protein up-regulated in carcinoma, membrane associated protein 17
2088	21695				ESTs, Weakly similar to CO1A_MOUSE CORONIN-LIKE PROTEIN P57 (CORONIN 1A) [M.musculus], coronin, actin binding protein 1A, coronin, actin binding protein 1B, coronin, actin binding protein 1C, coronin, actin-binding protein, 1A, hypothetical protein DKFZp7621166
		NM_130411	c,x		annexin VII, long form [H.sapiens], ESTs, Moderately similar to ANX4 MOUSE ANNEXIN IV [M.musculus], ZAP 36/annexin IV, annexin A4,
2089	21391	NM_130416 NM_130430	x,General		annexin A7
2090	20694 19818	NM_130430	General cc		
2090	18810	NM 130430	e,s		ATP synthase, H+ transporting, mitochondrial F1 complex, alpha subunit, isoform 1, ATP synthase, H+ transporting, mitochondrial F1 complex, alpha subunit, isoform 1, cardiac muscle, EST, Moderately similar to ATPA RAT ATP SYNTHASE ALPHA CHAIN, MITOCHONDRIAL PRECURSOR [R.norvegicus], expressed sequence AL022851, expressed sequence AL023067
2091	18293	NM_130433	q		
2092	25064	S45392	a,n		
2093	3244	S63519	u		
2094	25501	S63521	q		
2095	16248	S68135	h		EST, Weakly similar to ENPL_HUMAN ENDOPLASMIN PRECURSOR [H.sapiens], ESTs, Highly similar to HS9B RAT HEAT SHOCK PROTEIN HSP 90-BETA [R.norvegicus], ESTs, Weakly similar to ENPL_HUMAN ENDOPLASMIN PRECURSOR [H.sapiens], Homo sapiens mRNA; cDNA DKFZp564F053 (from clone DKFZp564F053), RIKEN cDNA 1810014B01 gene, RIKEN cDNA 2410002K23 gene, expressed sequence C81438, heat shock 90kD protein 1, beta, heat shock protein, 84 kDa 1, tumor rejection antigen (gp96)
2096	18647	S69316	q		1, tumor rejection antigen gp96
					ATP-binding cassette, sub-family D (ALD), member 4, ESTs, Highly similar to JC5604 ABC-transporting peroxisomal membrane protein 69 [H.sapiens], ESTs, Moderately similar to JC5604 ABC-transporting peroxisomal membrane protein 69
2097	24351	S74257	V		[H.sapiens]
2098	25066	S75280	d		L

TABLE	R HUMAN	Howorogae <u>v</u> v	INOTIATIONS		Atty. Docket No. 44921-5039W0 Doc. No. 1793897.1
839.ID No.	ldemiliter.	ConBank Ace./ Ref. Seq. (D No.	Model Gode	Homologous Cene Name 4: 1811	cansk reterið evogolomok
2099	1460 25539	S76054 S76742	j,l,m,x,y,General		DNA segment, Chr 15, Wayne State University 77, expressed, EST, Moderately similar to K2C8 RAT KERATIN, TYPE II CYTOSKELETAL 8 [R.norvegicus], EST, Weakly similar to I37982 Keratin 8 [H.sapiens], ESTs, Moderately similar to I37982 Keratin 8 [H.sapiens], Homo sapiens mRNA; cDNA DKFZp434C107 (from clone DKFZp434C107), RIKEN cDNA 1200016G03 gene, expressed sequence AL022697, expressed sequence AU019895, keratin 8, keratin complex 2, basic, gene 8
2100	16400	S76779			
2102	24469	S77858	n		EST, Weakly similar to MOHU6N myosin alkali light chain 6, nonmuscle form [H.sapiens], myosin light chain, alkali, nonmuscle, myosin, light polypeptide 6, alkali, smooth muscle and non-muscle
2103	25545	S77900	k,s		
2103	21583	S77900	k		EST, Weakly similar to MOHULP myosin regulatory light chain, placental [H.sapiens], ESTs, Moderately similar to MOHULP myosin regulatory light chain, placental [H.sapiens], myosin regulatory light chain, myosin, light polypeptide, regulatory, non-sarcomeric (20kD)
2104	10260	S81497	s	lipase A, lysosomal acid, cholesterol esterase (Wolman disease), lysosomal acid lipase 1	
2105	3609	S82579	k	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	Homo sapiens, Similar to histamine N- methyltransferase, clone MGC:14500 IMAGE:4249496, mRNA, complete cds, expressed sequence AI788969, histamine N-methyltransferase
2106	14959	U02506	a,q,General		EST, Moderately similar to GBLP_HUMAN GUANINE NUCLEOTIDE-BINDING PROTEIN BETA SUBUNIT-LIKE PROTEIN 12.3 [R.norvegicus], EST, Weakly similar to B33928 GTP-binding protein beta chain homolog [H.sapiens], ESTs, Weakly similar to A36986 activated protein kinase C receptor RACK1 - rat [R.norvegicus], Homo sapiens cDNA: FLJ21913 fis, clone HEP03888, Homo sapiens, Similar to guanine nucleotide binding protein (G protein), beta polypeptide 2-like 1, clone MGC:17239 IMAGE:4155303, mRNA, complete cds, expressed sequence AW544865, guanine nucleotide binding protein (G protein), beta polypeptide 2-like 1, guanine nucleotide binding protein, beta 2, related sequence 1

TABLES	REMUMAN	HOMOFOGRE VI	NOTATIONS		Any. Docket No. 44921-5039000 Doc. No. 1783397.1
Seg. ID No.	ldemiliter	ConBank Ace./ Ref. Seq. (D No.)	Model Gode &	Honologous Cens Nemo	Homologous Cluster Namo
2109 2110	2010 15462	U05675 U06230	b,x,bb		EST, Weakly similar to beta-fibrinogen precursor [H.sapiens], ESTs, Moderately similar to AF125176 1 angiopoietin-related protein-2 [M.musculus], ESTs, Weakly similar to FIBB RAT FIBRINOGEN BETA CHAIN PRECURSOR [R.norvegicus], expressed sequence AI256424, fibrinogen, B beta polypeptide
2112	1583	U07201	s,General	asparagine synthetase	
2113	627 809	U09229 U17035	h General	esperagine symmetase	ESTs, Highly similar to CDP_HUMAN CCAAT DISPLACEMENT PROTEIN [H.sapiens], Hepatocyte nuclear factor 6, Human chromosome 17q21 mRNA clone 1046:1-1, KIAA0293 protein, cut (Drosophila)-like 1, cut (Drosophila)-like 1 (CCAAT displacement protein), cut (Drosophila)-like 2, one cut domain, family member 1
	-	0.7.500	00.1014.	mini chromosome	
2115	16675	U17565	k,x,bb	maintenance deficient 6 (S. cerevisiae), minichromosome maintenance deficient (mis5, S. pombe) 6	mini chromosome maintenance deficient 6 (S. cerevisiae), minichromosome maintenance deficient (mis5, S. pombe) 6
2116	25587	U20110	r		
2117	90	U20796	r		Mus musculus, Similar to nuclear receptor subfamily 1, group D, member 1, clone MGC:6402 IMAGE:3585478, mRNA, complete cds, nuclear receptor subfamily 1, group D, member 2, thyroid hormone receptor alpha
2118 2119	25589 22196	U21718 U21719	h,aa h		
2120	17118	U25746	s		DEAD (aspartate-glutamate-alanine- aspartate) box polypeptide 5, ESTs, Moderately similar to A57514 RNA helicase HEL117 - rat [R.norvegicus], KIAA0801 gene product, RIKEN cDNA 2610007K22 gene, RIKEN cDNA 4921506D17 gene, RIKEN cDNA 9130430L19 gene, RNA helicase, expressed sequence Al325430
2121	1537	U27518	g,h,n		FOT Westless 2
2122	1558	U28504	ЬЬ		EST, Weakly similar to NPT1 RAT RENAL SODIUM-DEPENDENT PHOSPHATE TRANSPORT PROTEIN 1 [R.norvegicus], ESTs, Weakly similar to NPT1 MOUSE RENAL SODIUM-DEPENDENT PHOSPHATE TRANSPORT PROTEIN 1 [M.musculus], expressed sequence AW261723, solute carrier family 17 (sodium phosphate), member 1, solute carrier family 17 (sodium phosphate), member 3, solute carrier family 17 (sodium phosphate), member 4, solute carrier family 17 (sodium phosphate) member 4, solute carrier family 17 (sodium/hydrogen exchanger), member 1

Bone Bone	TABLES	DE CHUMMAN	HOMOFOGAE W	SKOTATOKI		Aiiy. Docket No. 44921-5089W0
BK protein, EST, Moderately simile St8989 EKX protein, Carp St Miles (R. norvegicus), Mus musculus St Mr RNA for BK protein, complete of strain RALBir, RIKEN cDNA C300008815 gene, synaptotagmin 5		n A control		m niân	Homologous Come	Dog. No. 179:597,1
2124	NO	icewayer.	Ker zed im No	Model Gode	Name	B/K protein, EST, Moderately similar to S68695 B/K protein - rat [R.norvegicus], Mus musculus B/K mRNA for B/K protein, complete cds,
2125				n		synaptotagmin 5
2126 25599 U34897 y						
Homo sapiens, clone MGC-4711 MMAGE:3334915, mRNA, complete				General		
2127 1394 U37099 h	2126	25599	U34897	ly		IMAGE:3534915, mRNA, complete cds, RAB23, member RAS oncogene family, RAB3A, member RAS oncogene family, RAB3C, member
ESTs, Highly similar to FGD1 - HUM PUTATIVE RHO/RAC GUANINE NUCLEOTIDE EXCHANGE FACTO (H.sapiens), ESTs, Weakly similar to B39898 phospholipase A2 (M.musculus), ESTs, Weakly similar to B39898 phospholipase A2 (M.musculus), ESTs, Weakly similar to FGD1 - HUMAN PUTATIVE RHO/RAC GUANINE NUCLEOTIDE EXCHANGE FACTOR (M.musculus), ESTs, Weakly similar to FGD1 - HUMAN PUTATIVE RHO/RAC GUANINE NUCLEOTIDE EXCHANGE FACTOR (H.sapiens), FGD1 family, member 3, RIKEN cDi 5830461L01 gene, faciogenital dysplasia homolog, faciogenital dysplasia homolog, faciogenital dysplasia homolog, faciogenital dysplasia homolog 2 (human) SETS, Weakly similar to AF167320 zinc finger protein ZFP113 (M.musculus), ESTs, Weakly similar to AF167320 zinc finger protein ZFP113 (M.musculus), ESTS, Weakly similar to ZF29 MOUSE ZIN FINGER PROTEIN 3] (H.sapiens), ESTS, Weakly similar to ZF29 MOUSE ZIN FINGER PROTEIN 13 (H.sapiens), ESTS, Weakly similar to ZF29 MOUSE ZIN FINGER PROTEIN 13 (H.sapiens), ESTS, Weakly similar to ZF29 MOUSE ZIN FINGER PROTEIN 13 (H.sapiens), ESTS, Weakly similar to ZF29 MOUSE ZIN FINGER PROTEIN 13 (H.sapiens), ESTS, Weakly similar to ZF29 MOUSE ZIN FINGER PROTEIN 13 (H.sapiens), ESTS, Weakly similar to ZF29 MOUSE ZIN FINGER PROTEIN 13 (H.sapiens), ESTS, Weakly similar to ZF29 MOUSE ZIN FINGER PROTEIN 13 (H.sapiens), ESTS, Weakly similar to ZF29 MOUSE ZIN FINGER PROTEIN 13 (H.sapiens), ESTS, Weakly similar to ZF29 MOUSE ZIN FINGER PROTEIN 2 (M.musculus), RIKEN cDNA 2310040101 gene, expressed sequence Al835008 (within H-2S), complement component 4 (within H-2S), complement component 4 (within H-2S), complement component C4A (H.sapiens) 2130	2127	1394	U37099	h		
PROTEIN 13 [H.sapiens], ESTs, Weakly similar to ZF29 MOUSE ZIN FINGER PROTEIN 29 [M.musculus, RIKEN cDNA 2310040101 gene, expressed sequence Al835008						ESTs, Highly similar to FGD1_HUMAN PUTATIVE RHO/RAC GUANINE NUCLEOTIDE EXCHANGE FACTOR [H.sapiens], ESTs, Weakly similar to B39898 phospholipase A2 [M.musculus], ESTs, Weakly similar to FGD1 MOUSE PUTATIVE RHO/RAC GUANINE NUCLEOTIDE EXCHANGE FACTOR [M.musculus], ESTs, Weakly similar to FGD1_HUMAN PUTATIVE RHO/RAC GUANINE NUCLEOTIDE EXCHANGE FACTOR [H.sapiens], FGD1 family, member 3, RIKEN cDNA 5830461L01 gene, faciogenital dysplasia (Aarskog-Scott syndrome), faciogenital dysplasia homolog, faciogenital dysplasia homolog 2 (human) ESTs, Weakly similar to AF167320 1 zinc finger protein ZFP113 [M.musculus], ESTs, Weakly similar to
Component Component Component Component Component Component Component Caspiens Component Caspiens Component Caspiens	2129	1623	U41164	ħ		PROTEIN 13 [H.sapiens], ESTs, Weakly similar to ZF29 MOUSE ZINC FINGER PROTEIN 29 [M.musculus], RIKEN cDNA 2310040101 gene,
2131 17886 U47315 S,Z	2130	15851	U42719	f.t.x.General	(within H-2S), complement	
2132 21654 U53184 i,t,General						The second and the second
2133						
expressed sequence AW146050, monocarboxylate transporter, solute carrier family 16 (monocarboxylic act transporters), member 3, solute carrifamily 16 (monocarboxylic acid transporters), member 3, solute carrifamily 16 (monocarboxylic acid transporters), member 7 2137 2153 U75404 b,cc,General D,cc,General 2133	1439					
2137 2153 U75404 b,cc,General protein (gravin) 12 2139 4956 U76714 j,y 2140 4477 U77829 I,m				bb		monocarboxylate transporter, solute carrier family 16 (monocarboxylic acid transporters), member 3, solute carrier family 16 (monocarboxylic acid
2139	2127	2153	1175404	h co Ganaral		
2140 4477 U77829 I,m expressed sequence C76683, putati					protein (gravin) 12	
\ expressed sequence C76683, putati						
					1	expressed sequence C76683, putative c-Myc-responsive

Sentition Sent	TABLE 8	e Humani	HOMOTOGAE VI	RYOTATOM		Aity. Docket No. 44921-5089W0
	Seq. ID No.	(den)iiler	Confork Acel Ref. See. ID No.	Model (2013)		
2143 23282		Marie and Communication of the				KIAA1683 protein, KIAA1802 protein, expressed sequence AA407558, lymphocyte antigen 64, polymerase (RNA) II (DNA directed) polypeptide A
2144 22005 U990725 h	2142	977	U89744	s		(S.cerevisiae) 5 homolog
2146 819						
State						
2147 818 X02291 e,j.z aldolase B, fructose-bisphosphate	2146	819	X02284	J,Z	Litata a O B is af	
2159					aldolase B, fructose-	
2150 20513 X05684 O,f Red blood cell, pyruvate kinase, liver and red blood cell, pyruvate kinase, liver and RBC						
2150 20513 X05684 O,r Kinase, liver and RBC	2149	16401	X04979	С		
25084 X06769 CC	2150	20513	X05684	0.5	red blood cell, pyruvate	
2152 672						
25675 X14181					· · · · · · · · · · · · · · · · · · ·	
EST, Moderately similar to RL1X_HUMAN 60S RIBOSOMAL PROTEIN L18A [H.sapiens], EST, Weakly similar to RL1X_HUMAN 60S RIBOSOMAL PROTEIN L18A, H.sapiens], EST, Weakly similar to RL1X_HUMAN 60S RIBOSOMAL PROTEIN L18A, cytosolic [H.sapiens], EST, estably similar to RL1X_HUMAN 60S RIBOSOMAL PROTEIN L18A, cytosolic [H.sapiens], ESTS, Highly similar to RL1X_HUMAN 60S RIBOSOMAL PROTEIN L18A [H.sapiens], ESTS, Highly similar to RL1X_HUMAN 60S RIBOSOMAL PROTEIN L26 [H.sapiens], ESTS, Highly similar to R13A, Highly						
BOS RIBŌSOMAL PROTEIN L26	2153	20810	X14181	n,q,w		RL1X_HUMAN 60S RIBOSOMAL PROTEIN L18A [H.sapiens], EST, Weakly similar to RL1X_HUMAN 60S RIBOSOMAL PROTEIN L18A [H.sapiens], EST, Weakly similar to S47353 ribosomal protein L18a, cytosolic [H.sapiens], ESTs, Highly similar to RL1X_HUMAN 60S RIBOSOMAL PROTEIN L18A [H.sapiens], RIKEN cDNA 2510019J09 gene, ribosomal protein L18a
EST, Weakly similar to 60S RIBOSOMAL PROTEIN L7A [M.musculus], ESTs, Highly similar to R5HU7A ribosomal protein L7a, cytosolic [H.sapiens], Homo sapiens rpL7a pseudogene, clone 3a, Humai DNA sequence from clone RP1- 189G13 on chromosome 20. Contair an RPL7A (60S ribosomal protein L7 (SURF3) pseudogene, an RPS4 (40 ribosomal protein S4) pseudogene, ESTs, STSs and GSSs, RIKEN cDN 4632404N19 gene, ribosomal protein L7a EST AI317031, EST, Weakly similar R3HU16 ribosomal protein S16, cytosolic [H.sapiens], expressed sequence AA420385, ribosomal protein S16 phospholipase A2, group				у		60S RIBOSOMAL PROTEIN L26 [H.sapiens], ESTs, Highly similar to S33713 ribosomal protein L26, cytosolic [H.sapiens], ESTs, Weakly similar to 60S RIBOSOMAL PROTEIN L26 [R.norvegicus], ribosomal protein L26, ribosomal protein L26
RIBOSOMAL PROTEIN L7A [M.musculus], ESTs, Highly similar to R5HU7A ribosomal protein L7a, cytosolic [H.sapiens], Homo sapiens rpL7a pseudogene, clone 3a, Human DNA sequence from clone RP1- 189G13 on chromosome 20. Contain an RPL7A (60S ribosomal protein L7 (SURF3) pseudogene, an RPS4 (40 ribosomal protein S4) pseudogene, ESTs, STSs and GSSs, RIKEN cDN 4632404N19 gene, ribosomal protein L7a 2155 19244 X15013 c.q,w EST Al317031, EST, Weakly similar R3HU16 ribosomal protein S16, cytosolic [H.sapiens], expressed sequence AA420385, ribosomal protein S16	2155	25679	X15013	q		
R3HU16 ribosomal protein S16, cytosolic [H.sapiens], expressed sequence AA420385, ribosomal protein S16 2156	2155	19244	X15013	c,q,w		RIBOSOMAL PROTEIN L7A [M.musculus], ESTs, Highly similar to R5HU7A ribosomal protein L7a, cytosolic [H.sapiens], Homo sapiens rpL7a pseudogene, clone 3a, Human DNA sequence from clone RP1-189G13 on chromosome 20. Contains an RPL7A (60S ribosomal protein L7A) (SURF3) pseudogene, an RPS4 (40S ribosomal protein S4) pseudogene, ESTs, STSs and GSSs, RIKEN cDNA 4632404N19 gene, ribosomal protein L7a
phospholipase A2, group			;			R3HU16 ribosomal protein S16, cytosolic [H.sapiens], expressed
	2156	15626	X17665	а		protein S16
	2157	1802	X51520	+	IIA (platelets, synovial	
2158 25686 X51536 bb				hh	liuiu)	

TABLE 8	B HUMAN	HOMOLOGUE AN	EXIOTATION		Ally. Docket No. 44921-51139WO Doc. No. 1793397.1
Seq. [D No. 4	ldeniiier	Cenbenk Ace./ Ref. Seq. ID No.	Model Code	Homologous Como Nemo	Homologous Gluster Name
		V			EST, Moderately similar to R3RT3 ribosomal protein S3 - rat [R.norvegicus], EST, Weakly similar to R3RT3 ribosomal protein S3 - rat
2158	10819	X51536	aa,bb		[R.norvegicus], hypothetical protein FLJ23059, ribosomal protein S3 EST, Weakly similar to 60S
					RIBOSOMAL PROTEIN L9 [R.norvegicus], EST, Weakly similar to S42106 ribosomal protein L9 homolog [H.sapiens], ESTs, Highly similar to S42106 ribosomal protein L9 homolog [H.sapiens], ESTs, Moderately similar to 60S RIBOSOMAL PROTEIN L9 [R.norvegicus], RIKEN cDNA 4930401B11 gene, ribosomal protein
2159	18250	X51706	a,q,w	ribosomal protein L9	L9 EST, Moderately similar to 40S
					RIBOSOMAL PROTEIN S19 [R.norvegicus], EST, Weakly similar to RS19_HUMAN 40S RIBOSOMAL PROTEIN S19 [H.sapiens], ribosomal
2160	20872	X51707	а	ribosomal protein S19	protein S19 myxovirus (influenza virus) resistance
2464	540	VE0744			1, myxovirus (influenza) resistance 1, homolog of murine (interferon-
2161 2162	516 25689	X52711 X52815	g		inducible protein p78)
2163	20427	X53378	w		ESTs, Highly similar to RS13_HUMAN 40S RIBOSOMAL PROTEIN S13 [H.sapiens], ESTs, Moderately similar to RS13_HUMAN 40S RIBOSOMAL PROTEIN S13 [H.sapiens]
2164	18606	X53504	General		EST, Moderately similar to S35531 ribosomal protein L12, cytosolic [H.sapiens], EST, Weakly similar to 60S RIBOSOMAL PROTEIN L12 [R.norvegicus], hypothetical protein, ribosomal protein L12
2165	1463	X54467	d,u,General		EST, Weakly similar to 60S ACIDIC
					RIBOSOMAL PROTEIN P2 [R.norvegicus], EST, Weakly similar to R6HUP2 acidic ribosomal protein P2, cytosolic [H.sapiens], ESTs, Highly similar to 60S ACIDIC RIBOSOMAL PROTEIN P2 [R.norvegicus], ESTs, Highly similar to MTJ1 MOUSE DNAJ PROTEIN HOMOLOG MTJ1 [M.musculus], Human DNA sequence from clone RP3-408B20 on chromosome 6 Contains ESTs, STSs and GSSs. Contains a gene and two pseudogenes for novel 7 transmembrane receptors (olfactory family) and a gene for a novel protein similar to 60S acidic ribosomal protein
2166	24577	X55153	a,v		P2 (RPLP2), RIKEN cDNA 2700049122 gene, ribosomal protein, large P2, ribosomal protein, large, P1

TABLE		HOMOLOGUEAN	NOTATIONS		//Aity. Docket No. 44924-5039V/ Doc. No. 1793397.
509. (ID No.	lentifier	Coneank Aced Ref. Seq. ID No.		Honologous Cane	Homologous Cluster Name
Baseline and out-of-service				And the control of th	EST, Highly similar to A40043 notch protein homolog TAN-1 precursor [H.sapiens], EST, Weakly similar to A40043 notch protein homolog TAN-1 precursor [H.sapiens], ESTs, Weakly similar to NEUROGENIC LOCUS NOTCH HOMOLOG PROTEIN 1 PRECURSOR [M.musculus], Homo sapiens mRNA; cDNA DKFZp761G02121 (from clone DKFZp761G02121); partial cds, Notch (Drosophila) homolog 1 (translocationassociated), Notch (Drosophila)
				Notch (Drosophila) homolog 1 (translocation- associated), Notch gene	homolog 2, Notch (Drosophila) homolog 3, Notch 3, Notch gene homolog 1, (Drosophila), Notch gene
2167	10344	X57405	j,m	homolog 1, (Drosophila)	homolog 3, (Drosophila), jagged 1 EST, Weakly similar to S30393 ribosomal protein S18, cytosolic [H.sapiens], ESTs, Highly similar to S30393 ribosomal protein S18,
2168	15106	X57529	g,n,q		cytosolic [H.sapiens], ribosomal protein S18
2169	5667	X58200	q,bb	ribosomal protein L23	protein 010
2169	18611	X58200	a,v	ribosomal protein L29	RIBOSOMAL PROTEIN L29 [R.norvegicus], EST, Weakly similar to S65784 ribosomal protein L29, cytosolic [H.sapiens], ESTs, Highly similar to S65784 ribosomal protein L29, cytosolic [H.sapiens], ESTs, Weakly similar to 60S RIBOSOMAL PROTEIN L29 [M.musculus], Human DNA sequence from clone RP4-595K12 on chromosome 1p31.2-31.3 Contains a pseudogene similar to 60S RPL29 (ribosomal protein L29 (cell surface heparin binding protein HIP)), a chromosome 1 specific mRNA (KIAA0499), a novel mRNA (KIAA0433), ESTs, STSs, GSSs and a CpG Island, ribosomal protein L29
					EST, Moderately similar to RL17_HUMAN 60S RIBOSOMAL PROTEIN L17 [H.sapiens], EST, Weakly similar to RL17 RAT 60S RIBOSOMAL PROTEIN L17 [R.norvegicus], EST, Weakly similar to RL17_HUMAN 60S RIBOSOMAL PROTEIN L17 [H.sapiens], ESTs, Weakly similar to R5HU22 ribosomal
2170	17175	X58389	w		protein L17, cytosolic [H.sapiens] EST, Moderately similar to 40S RIBOSOMAL PROTEIN S5 [R.norvegicus], ESTs, Weakly similar to S55916 ribosomal protein S5,
2171	25702	X58465	w	ribosomal protein S5	cytosolic [H.sapiens], ribosomal protein S5 EST, Moderately similar to 40S RIBOSOMAL PROTEIN S5 [R.norvegicus], ESTs, Weakly similar to 555916 ribosomal protein S5, cytosolic [H.sapiens], ribosomal
2171	10109	X58465	c,q	ribosomal protein S5	protein S5
2172	25705	X59375	c,i,aa,General		

TABLES	DE CHUMAN	HOMOTOGAE VA	EXPORTATIONS		Ally. Docket No. 44921-5039W0
Scq. (D		ConBank Acc./		Homologous Gana	
	ldentifier			Name :	Homologovs Cluster Name
2173	25709	X59737	lu	creatine kinase, mitochondrial 1 (ubiquitous), creatine kinase, mitochondrial 1, ubiquitous	
2173	20703	7,557.67	<u> </u>	abiquitous	ESTs, Moderately similar to dJ63G5.3
2174	18354	X59859	General	decorin	[H.sapiens], RIKEN cDNA 1700034K16 gene, RIKEN cDNA 5530600M07 gene, decorin
					ESTs, Moderately similar to dJ63G5.3 [H.sapiens], RIKEN cDNA 1700034K16 gene, RIKEN cDNA
2174	18355	X59859	t	decorin	5530600M07 gene, decorin ESTs, Moderately similar to S17182
2175	21657	X61381	General		interferon-induced protein 1-8U [H.sapiens], ESTs, Weakly similar to putative haemopoietic membrane protein [M.musculus], Human DNA sequence from clone RP4-781L3 on chromosome 1p34.3-36.11 Contains a pseudogene similar to IFITM3 (interferon inducedntransmembrane protein 3 (1-8U)), STSs and GSSs, RIKEN cDNA 1110004C05 gene, interferon induced transmembrane protein 3 (1-8U), interferon induced transmembrane protein 3-like, interferon-inducible protein 16
2176	25718	X62145	bb,General		
2176	15875	X62145	a,q,v		EST, Highly similar to 60S RIBOSOMAL PROTEIN L8 [R.norvegicus], EST, Moderately similar to 60S RIBOSOMAL PROTEIN L8 [R.norvegicus], EST, Weakly similar to JN0923 ribosomal protein L8, cytosolic [H.sapiens], ESTs, Highly similar to 60S RIBOSOMAL PROTEIN L8 [R.norvegicus], ESTs, Highly similar to RL8_HUMAN 60S RIBOSOMAL PROTEIN L [M.musculus], ESTs, Moderately similar to RL8_HUMAN 60S RIBOSOMAL PROTEIN L [M.musculus], expressed sequence AL024098, ribosomal protein L8
2177	13646	X62166	bb		EST, Weakly similar to 60S RIBOSOMAL PROTEIN L3 [R.norvegicus], EST, Weakly similar to I84501 ribosomal protein L3 [H.sapiens], ESTs, Moderately similar to 60S RIBOSOMAL PROTEIN L3 [R.norvegicus], ESTs, Moderately similar to 184501 ribosomal protein L3 [H.sapiens], ESTs, Weakly similar to RL3 MOUSE 60S RIBOSOMAL PROTEIN L3 [M.musculus], RIKEN cDNA 1110057H16 gene, ribosomal protein L3, ribosomal protein L3-like
2178	25721	X62325	р		
2179	16012	X62875	m,s,z		high mobility group AT-hook 1, high- mobility group (nonhistone chromosomal) protein isoforms I and Y
					
2180 2181	25730 25089	X63369 X63594	cc General		· · · · · · · · · · · · · · · · · · ·

TABLES		HOWOROGALE YA	NOTATIONS:	E MANA E E E E	Alty, Docket No. 44921-5089W0
569. ID No.	ldentifier	Cenbenk Ace/ Ref. Seq. ID No.	Model Code	Homologous Cene Name	Doc. No. 1793397.1 Homologous Gluster Name
	Ballion Co. An Eastern Household Co.	Section 1 Access to the Control of t		The state of the s	EST, Highly similar to 60S RIBOSOMAL PROTEIN L23A [R.norvegicus], ESTs, Highly similar to RL2B_HUMAN 60S RIBOSOMAL PROTEIN L23A [H.sapiens], ribosomal
2182	20844	X65228	n,w		protein L23a
2183 2184	20879 25736	X65296 X68782	j.y c		EST, Weakly similar to JC5408 carboxylesterase [H.sapiens], ESTs, Moderately similar to ES22 MOUSE LIVER CARBOXYLESTERASE 22 PRECURSOR [M.musculus], T-complex expressed gene 5, carboxylesterase 1, carboxylesterase 1 (monocyte/macrophage serine esterase 1), carboxylesterase 3, carboxylesterase 3 (brain)
2184	25/36	X68782	С		EST Highly similar to CA13 HUMAN
2185	16426	X70369 X70706	с	collagen, type III, alpha 1 (Ehlers-Danlos syndrome type IV, autosomal dominant), procollagen, type III, alpha 1	EST, Highly similar to CA13_HUMAN COLLAGEN ALPHA 1(III) CHAIN PRECURSOR [H.sapiens], EST, Moderately similar to CA13_HUMAN COLLAGEN ALPHA 1(III) CHAIN PRECURSOR [H.sapiens], EST, Weakly similar to CGHU7L collagen alpha 1(III) chain precursor [H.sapiens], ESTs, Highly similar to CA21_HUMAN COLLAGEN ALPHA 2(I) CHAIN PRECURSOR [H.sapiens], collagen type V, alpha 2, collagen, type III, alpha 1 (Ehlers-Danlos syndrome type IV, autosomal dominant), macrophage receptor with collagenous structure, procollagen, type III, alpha 1 ESTs, Highly similar to A34789 T-plastin [H.sapiens], ESTs, Highly similar to PLSI_HUMAN I-PLASTIN [H.sapiens], expressed sequence Al115446, expressed sequence Al427122, expressed sequence AL024105, plastin 2, L
					B-cell CLL/lymphoma 1, EST, Moderately similar to CGD1 RAT G1/S- SPECIFIC CYCLIN D1 [R.norvegicus], ESTs, Weakly similar to 1709356A cyclin PRAD1 [H.sapiens], cyclin D1, cyclin D1 (PRAD1: parathyroid adenomatosis 1), expressed sequence
2187	24232	X75207	С		Al327039
2188 2189	16272 25741	X76456 X76489	n,p		ESTs, Highly similar to alpha-albumin protein [M.musculus], Mus musculus mRNA for alpha-albumin protein, afamin
2109	20141	N 0403	<u> </u>		ESTs, Weakly similar to DAHUA1
2190	23302	X78949	h		procollagen-proline dioxygenase [H.sapiens], expressed sequence Al853847, expressed sequence C76437, procollagen-proline, 2-oxoglutarate 4-dioxygenase (proline 4-hydroxylase), alpha 1 polypeptide, procollagen-proline, 2-oxoglutarate 4-dioxygenase (proline 4-hydroxylase), alpha polypeptide I, procollagen-proline, 2-oxoglutarate 4-dioxygenase (proline 4-hydroxylase), alpha polypeptide II

TABUE 8	KIMMUN)	HOMOLOGUEAN		The module of the state of the	Ally, Docket No. 44921-5039W0 Doc. No. 1798397.1
899. ID No.	ldentifier	Coneank Ace <i>l</i> Rof. Soq. ID No.	Model Code	Homologous Cene Name	Homologous Cluster Name
2191	25747	X81448	General		
2192	24115	X81449	u		EST, Weakly similar to KERATIN, TYPE I CYTOSKELETAL 19 [M.musculus], ESTs, Moderately similar to K1CJ_HUMAN KERATIN, TYPE I CYTOSKELETAL 10 [H.sapiens], ESTs, Weakly similar to S30433 keratin 17, type I, cytoskeletal [H.sapiens], Homo sapiens mRNA for keratin 19, partial cds, isolate:K19-141 keratin 19, keratin complex 1, acidic, gene 19, type I intermediate filament cytokeratin
2193	25754	X89696	g		
2194	25097	X90642	y,z		
2195	12978	X96437	cc,General		immediate early response 3
2197	4594	Y07704	С		
2199	15986 20890	Y09945 Y13275	g,p,General bb,General		EST, Weakly similar to OCN2 MOUSE ORGANIC CATION/CARNITINE TRANSPORTER 2 [M.musculus], ESTs, Highly similar to OCN2_HUMAN ORGANIC CATION/CARNITINE TRANSPORTER 2 [H.sapiens], ESTs, Weakly similar to JE0346 high-affinity carntine transporter, CT1 - rat [R.norvegicus], ion transporter protein, solute carrier family 22 (organic cation transporter), member 1, solute carrier family 22 (organic cation transporter), member 4, solute carrier family 22 (organic cation transporter), member 5, solute carrier family 22 (organic cation transporter), member 9 CD9 antigen, RIKEN cDNA 6330415F13 gene, RIKEN cDNA B230119D02 gene, expressed sequence C76990, transmembrane 4 superfamily member 3 ESTs, Weakly similar to DAD1_HUMAN DEFENDER AGAINST
2201	21914	Y13336	d		CELL DEATH 1 [R.norvegicus], defender against cell death 1, expressed sequence Al323713
2202	406	Z11995	o,General	İ	1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1
2203 2204	18352 17481	Z12298 Z49761	t k	decorin	ESTs, Moderately similar to dJ63G5.3 [H.sapiens], RIKEN cDNA 1700034K16 gene, RIKEN cDNA 5530600M07 gene, decorin
2205	8664	Z75029	r,v		ESTs, Highly similar to T17342 hypothetical protein DKFZp586K1924.1 [H.sapiens], ESTs, Moderately similar to T17342 hypothetical protein DKFZp586K1924.1 [H.sapiens]
2206	2459	AA964755	cc		
2207	23830	AA956638	aa		
2208	6100	X73524	x		

TABLE:	B. HUMAN	HONOFOGNE W	RIOTEATIONS		Ally, Docket No. 44921-5089W0 Doc, No. 1798397.1
80g. ID No.	ldeniliter	Cenbenk Acel Ref. Seq. 10 No.	Model Gode	Homologous Cene Neme	Homologous Cluster Name
2209	439	Z22607	w	bone morphogenetic protein 4	bone morphogenetic protein 15, bone morphogenetic protein 4, endometrial bleeding associated factor, endometrial bleeding associated factor (left-right determination, factor A; transforming growth factor beta superfamily), growth differentiation factor 2, growth differentiation factor 5, hypothetical protein FLJ10314 ESTs, Highly similar to T17342
2210	8665	Al071965	v		hypothetical protein DKFZp586K1924.1 [H.sapiens], ESTs, Moderately similar to T17342 hypothetical protein DKFZp586K1924.1 [H.sapiens]
2211	155	U32681	t	selenium glutathione peroxidase, acidic calcium- independent	CD163 antigen, ESTs, Highly similar to I38005 M130 antigen precursor, splice form 4 [H.sapiens], KIAA1822 protein, apoptosis inhibitory 6, crp-ductin, deleted in malignant brain tumors 1, lectin, galactoside-binding, soluble, 3 binding protein, macrophage scavenger receptor 2, peptidylprolyl isomerase C-associated protein ESTs, Moderately similar to AOP2_HUMAN ANTIOXIDANT PROTEIN 2 [H.sapiens], anti-oxidant protein 2 (non-selenium glutathione peroxidase, acidic calcium-
2212	19252	AA892041	s	phospholipase A2), peroxiredoxin 5	independent phospholipase A2), peroxiredoxin 5
2213	15582	AI232320	q		
2214	17541	M26125	n	epoxide hydrolase 1, microsomal, epoxide hydrolase 1, microsomal (xenobiotic)	EST, Moderately similar to HYEP_HUMAN EPOXIDE HYDROLASE 1 [H.sapiens], ESTs, Highly similar to HYEP_HUMAN EPOXIDE HYDROLASE 1 [H.sapiens], epoxide hydrolase 1, microsomal (xenobiotic)
2215	18609	M30689	i		
2216	6262	AI177125	g		
2217	23859	AI072161 H32189	e	glutathione S-transferase M2 (muscle), glutathione S- transferase, mu 2	ESTs, Moderately similar to GLUTATHIONE S-TRANSFERASE YB1 [R.norvegicus], glutathione S- transferase M1, glutathione S- transferase, mu 1
2220	2572	AI177143	b		
2221	25419	M22922	а		

TABLE 4: GODE KEY	Aity. Docket No. 44921-5089W0 Doc. No. 1798397.1			
	Time(brs)	Code		
GENERAL TOXICITY		General		
ACYCLOVIR	24, 168	а		
ACYCLOVIR	6	b		
ADR	120, 168	С		
AY	360	d		
BEA	6, 24	е		
CAPTOPRIL	336	f		
CARBOPLATIN	6	g		
CEPHALORIDINE	6, 24	h		
CIDOFOVIR	120	i		
CISPANcombined	6, 24	j		
CISPLATIN	168	k		
CISPLATIN	6, 24	1		
CISPLATIN	6, 24, 168	m		
CITRININ	6, 24	n		
COLCHICINE	6, 24, 48	0		
CYCLOPHOSPHAMIDE	6	р		
DIFLUNISAL	24	q		
HYDRALAZINE	6	r		
IFOSFAMIDE	6, 24, 48, 144	S		
INDOMETHACIN	48, 72	t		
LITHIUMCHLORIDE	120	u		
MERCURICCHLORIDE	3, 6, 24	V		
PAMIDRONATE	24	w		
PAN	168	x		
PAN	6, 24	у		
PAN	6, 24, 168	Z		
SEMUSTINE	168	aa		
SULFADIAZINE	24	bb		
SULFADIAZINE	3, 6	СС		

		-370	0-			
* *	GENERAL			ket No. 44 Doc. N	1921-5039VVC 10. 1798397.1	
ldendifier		NonToxSD	ToxMean	ToxSD	I was a second of the second o	
12979	326.05	98.48	729.13	345.15	83.46	
23314	-7.57	63.50	480.54	528.59	83.43	
5461	161.65	48.19	368.42	210.09	82.70	
9583	40.60	17.67	158.01	127.01	82.56	
16982	59.51	36.53	506.74	577.57	81.34	
1809	5.28	13.85	191.53	265.23	81.21	
19184	59.98	26.06	191.02	128.18	81.04	
24200	382.07	78.91	618.97	199.18	80.18	
15003	13.42	14.74	154.63	217.73	79.83	
2629	18.92	8.93	58.17	43.50	79.72	
22321	82.69	25.99	192.94	130.47	79.58	
15301	20.84	20.65	124.72	132.03	79.47	
15032	280.18	50.93	183.93	78.32	79.33	
7489	89.81	28.30	47.02	25.94	79.28	
2242	2431.04	453.24	1658.86	711.96	79.26	
3050	77.80	26.29	166.37	91.49	79.22	
22681	170.38	56.88	497.37	377.15	79.20	
24042	4.31	9.28	112.52	198.98	79.06	
14425	191.57	51.38	315.84	117.32	79.05	
15300	104.10	41.29	305.26	261.16	79.01	
23651	487.82	171.51	1473.41	1227.36	78.44	
15964	1274.63	262.79	824.24	343.96	78.27	
16312	44.81	17.42	107.40	60.46	78.21	
16168	305.21	53.33	588.32	425.68	78.12	
5384	28.68	24.27	102.33	70.63	78.08	
12978	92.29	28.22	178.30	79.32	77.90	
21654	332.77	53.62	518.29	196.63	77.78	
3874	934.64	159.07	672.84	188.60	77.60	
3049	176.71	56.69	334.05	163.05	77.39	
16314	33.02	22.70	103.37	64.81	77.39	
23299	345.71	73.26	514.49	166.24	77.35	
9166	13.09	10.08	40.42	28.18	77.33	
14763	1.59	39.67	228.03	268.46	77.30	
4479	133.60	62.59	233.57	86.27	77.27	
15928	142.69	34.11	244.10	90.89	77.08	
3941	229.39	59.58	325.62	88.12	76.99	
28	540.29	128.18	342.30	188.50	76.94	
14929	687.79	150.66	1358.82	807.54	76.87	
22885	1229.56	342.00	1987.11	703.59	76.84	
22765	15.12	10.37	52.20	40.27	76.68	
19040	158.12	34.83	333.27	212.49	76.68	
21239	98.57	31.29	190.62	93.31	76.57	
2555	83.76	25.31	159.42	79.77	76.51	
15051	558.01	156.42	953.70	424.38	76.38	
22569	701.48	148.25	468.05	181.60	76.33	
15299	73.40	22.15	164.49	121.44	76.17	
20116	1.36	12.44	53.19	60.94	76.12	
7299	141.75	59.67	363.99	279.53	75.86	
11618	445.83	108.35	290.61	162.81	75.81	
23868	113.35	64.90	514.43	654.54	75.65	
312	164.76	29.18	118.06	38.83	75.63	
23166	116.47	37.76	215.89	104.73	75.55	
19723	63.30	24.96	147.23	109.39	75.53	
2161	10.89	22.79	40.17	28.34	75.53	
22592	195.01	88.25	453.16	291.73	75.50	
21683	27.66	14.45	65.00	37.84	75.44	

TABLE 5:	GENERAL *		Aity. Docket No. 44 921-5039 W0 Doc. No. 1793397.1			
nelfiffmelbl	NonToxMean:	เลราจาโกคที่	ToxWeam		LDAScore	
7540	135.61	39.51	269.18	167.73	75.42	
3121	1387.78	270.45	970.56	410.36	75.23	
17325	37.68	26.79	190.12	198.72	75.23	
4049	8.16	14.26	100.05	143.11	75.21	
24219	294.84	62.68	415.01	115.05	75.14	
7101	266.61	65.47	1024.56	1594.33	75.14	
21462	246.43	47.45	320.88	67.53	75.11	
21458	203.89	61.93	345.78	139.58	75.01	
1460	178.45	37.93	302.50	171.15	74.92	
23957	57.89	29.03	120.87	65.67	74.89	
12921	92.69	31.09	174.99	86.55	74.88	
24237	46.61	22.12	105.68	71.47	74.84	
20830	482.35	118.55	717.12	282.77	74.80	
14185	181.85	52.63	307.94	156.98	74.74	
3091	821.34	154.51	616.81	215.94	74.73	
6046	221.77	64.88	141.08	71.40	74.73	
10818	509.63	159.26	301.00	207.18	74.63	
18906	270.92	73.69	171.86	79.69	74.60	
17361	160.53	54.00	92.22	55.73	74.60	
574	297.07	48.04	543.78	340.56	74.58	
1529	305.18	51.69	224.24	69.41	74.56	
20161	30.38	21.73	80.33	56.50	74.54	
22152	-0.67	16.93	51.48	64.42	74.54	
21391	183.11	55.68	391.43	249.63	74.34	
20056	319.93	46.22	248.41	81.57	74.34	
5711	402.83	114.80	268.76	100.84	74.34	
16169	126.24	66.19	419.86	462.11	74.32	
7196	160.17	37.22	297.55	159.08	74.29	
13634	754.99	133.63	1123.83	475.68	74.29	
10659	111.21	38.66	239.07	166.99	74.28	
15089	162.94	57.42	271.19	109.36	74.26	
2628	8.82	14.55	37.93	39.15	74.22	
1521	7.84	37.91	80.00	74.26	74.20	
17524	1225.79	235.19	927.05	284.85	74.20	
14677	64.57	22.56	114.45	54.63	74.04	
17357	284.47	66.73	189.90	103.16	73.97	
15382	79.92	58.73	367.62	435.18	73.94	
1141	226.12	47.74	315.88		73.94	
3995	643.39	131.36	476.67	160.12	73.88	
6804	1354.29	374.19	830.60	415.24	73.82	
20694	1004.30	200.48	773.76	246.63	73.81	
8477	493.77	117.87	724.70	226.98	73.80	
13332	440.18	81.35	336.07	113.36	73.79	
2912	2775.27	619.46	2040.05	590.49	73.79	
8143	30.96	28.78	105.91	87.78	73.79	
8639	351.67	79.66	468.70	121.28	73.78	
354	191.57	43.15	335.31	175.98	73.77	
2702	261.10	53.61	363.77	115.07	73.74	
13411	857.90	304.76	508.64	242.77	73.71	
23261	1651.29	316.95	1185.07	374.72	73.69	
16775	1092.58	348.47	732.47	398.39	73.66	
10016	191.31	45.71	298.01	134.57	73.59	
353	154.16	43.72	275.53	149.36	73.59	
		56.55	318.50	126.31	73.55	
5295	199.26	เอต.ออ	เอ เด.อบ	1 1 2 0 3 1	17 3.00	

TABLE 5:	CENERAL		Aliy. Docket No. 4 Doc. [1921 -5009WG Vo. 1798397.1	
ldendifter:	MonToxMean	NonToxSD	ToxiMean	ToxsD	LDAScore	
20458	377.19	83.99	283.38	95.28	73.52	
23869	24.11	23.82	132.98	190.30	73.49	
20848	474.69	80.77	708.43	261.80	73.46	
9067	645.45	126.00	841.67	201.55	73.45	
923	10.66	7.41	32.76	27.77	73.40	
4291	317.92	87.25	202.30	99.83	73.38	
18529	184.43	47.69	306.78	139.66	73.35	
22626	66.24	25.65	191.03	161.08	73.32	
3823	488.87	101.91	709.19	233.23	73.30	
15663	179.56	40.22	269.94	97.24	73.29	
22929	927.47	283.23	548.56	310.91	73.27	
373	20.21	24.84	107.75	123.23	73.25	
4952	97.10	31.18	160.25	66.14	73.23	
2905	221.02	70.67	379.00	165.35	73.23	
7127	301.01	84.77	195.82	87.14	73.22	
20035	157.82	53.67	318.27	196.89	73.21	
14424	40.56	38.11	216.48	294.45	73.19	
1501	46.55	22.43	125.28	104.55	73.19	
811	268.35	42.34	201.54	69.48	73.19	
				411.90	73.12	
3610	1272.79	264.85 41.36	879.10			
9053	249.79		192.96	58.86	73.09	
23538	85.95	40.73	190.24	133.78	73.07	
18337	1699.33	315.35	1236.46	390.84	73.06	
15002	119.96	26.12	252.22	220.20	73.06	
21147	365.33	62.56	285.83	76.45	73.05	
8721	208.57	63.43	132.86	65.90	73.01	
1462	364.98	75.98	659.63	429.84	73.01	
11483	54.82	19.42	118.04	81.85	72.90	
2348	545.81	197.73	349.07	201.06	72.86	
1564	6.71	7.78	209.28	375.95	72.86	
12467	60.46	20.47	103.84	48.93	72.85	
6638	104.00	28.03	76.61	32.45	72.83	
19031	50.08	27.00	116.79	84.17	72.83	
1246	98.28	29.26	60.78	37.90	72.82	
23872	30.55	27.67	146.46	207.24	72.80	
19678	121.25	48.99	54.47	69.08	72.78	
23512	1086.33	216.78	839.09	240.27	72.77	
6321	458.69	127.59	644.50	219.13	72.75	
22596	57.04	17.25	81.28	26.16	72.75	
24431	50.52	15.50	151.16	210.56	72.73	
15110	663.97	145.61	479.15	149.17	72.73	
15892	12.32	15.49	44.12	34.64	72.71	
14458	29.92	20.00	71.32	39.77	72.67	
6641	402.83	66.65	309.63	82.93	72.53	
1422	315.77	80.70	215.58	83.38	72.53	
21443	90.59	31.63	155.73	92.87	72.53	
8829	264.33	62.24	350.44	106.21	72.51	
21632	27.74	33.70	87.38	68.91	72.48	
24388	173.36	44.15	275.21	119.72	72.47	
15851	171.96	68.96	332.47	267.49	72.38	
15042	55.43	27.40	117.95	81.96	72.32	
17908	49.38	21.99	125.62	121.11	72.32	
17908 15618			124.19		72.27	
21318	90.24 41.50	22.13 23.94	72.71	32.66 32.13	72.27	
	(4 L 3U	17.3 94	1///	1.37 1.5	11///	

TABLE 5:	CENERAL .	Karan Araban Karan	Ally. Doc	kot No. 44	1921- 5009 000
				T	<u>lo. 179339</u> 7.1
ldendifer	MOONTOXME	NonToxSD	ToxiMean	Toxsd	LDAScore
1727	30.19	25.05	113.76	121.46	72.19
23202	169.96	36.51	126.76	36.17	72.16
22248	216.64	78.55	393.59	188.26	72.15
22612	487.17	110.76	359.48	126.14	72.14
17734	106.27	34.17	214.91	182.70	72.12
19235	1112.95	271.69	810.00	308.94	72.06
13618	96.22	26.36	137.92	45.79	72.04
19525	23.64	12.50	46.38	27.18	72.03
4584	76.41	22.50	109.66	36.31	71.98
22197	112.68	34.33	178.88	75.61	71.98
24762	1064.07	310.38	755.53	273.04	71.97
10985	1189.01	218.09	885.13	290.39	71.96
3145	466.94	133.08	330.71	144.92	71.93
20828	278.75	83.71	482.02	272.83	71.89
2395	172.58	47.67	175.81	100.28	71.86
13609	252.08	52.43	186.04	64.47	71.86
21339	29.51	15.60	53.28	32.70	71.78
3079	27.52	22.01	67.24	49.93	71.78
16321	219.91	42.72	273.17	55.96	71.77
4944	100.32	33.19	177.16	86.89	71.77
24568	162.64	51.57	105.85	46.68	71.75
3875	539.59	120.20	389.50	146.06	71.75
6382	117.04	33.24	171.65	55.00	71.73
3959	329.28	86.36	451.00	151.79	71.73
8795	14.06	10.71	26.30	13.30	71.73
					1
17477	102.32	22.97	156.34	68.17	71.71
7700	74.32	20.26	138.95	92.87	71.68
17550	1380.51	319.12	1028.40	363.64	71.63
410	1155.16	215.31	918.14	252.51	71.63
17682	706.42	143.23	505.72	215.80	71.63
5897	20.54	16.83	45.41	23.66	71.63
4661	288.80	58.15	408.62	120.87	71.59
16521	266.79	60.39	378.60	127.02	71.59
13610	371.99	53.12	283.18	88.22	71.55
22554	565.54	122.08	428.04	154.62	71.49
11910	25.51	35.83	-5.43	31.38	71.47
15588	-4.65	25.14	33.98	37.77	71.42
5601	1014.34	210.01	731.23	341.39	71.42
5780	-23.15	27.07	32.19	62.08	71.38
21546	-79.60	43.55	44.91	170.38	71.38
15039	285.29	65.31	206.99	101.12	71.38
18300	483.67	133.71	307.50	148.77	71.38
14970	215.37	35.86	161.84	54.69	71.32
13151	635.29	173.43	1103.53	626.81	71.31
7197	180.11	57.15	296.88	139.25	71.28
21238	-24.44	33.42	22.20	43.34	71.27
25090	74.42	33.18	128.49	62.02	71.24
18564	225.23	44.60	181.56	58.17	71.23
1409	441.80	78.21	357.51	91.06	71.20
7903	526.91	184.33	331.16	186.75	71.18
24109	231.59	87.11	172.36	143.80	71.18
3416	108.18	41.02	234.59	190.53	71.17
2250	1462.04	256.17	1165.90	304.84	71.12
5867	157.83	35.87	210.76	63.63	71.11
17771	710.83	199.94	1089.58	460.00	71.09

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TABLE 5:	General :		AMy. Doc	14: Doc. 14 Doc. 14	1921-5039\WQ 10. 1793397.1
ldentifer	NonToxMean	NonToxSD	ToxMean	ToxSD	LDAScore
5494	62.28	23.39	104.89	53.79	71.08
18269	729.55	160.51	578.31	143.90	71.07
14996	459.09	95.27	332.24	132.64	71.03
24617	27.48	28.57	-0.27	26.14	71.02
23195	323.13	74.62	242.16	107.55	71.01
22656	113.29	44.19	184.20	80.34	70.99
8728	114.64	32.58	163.70	57.48	70.99
14664	56.90	24.89	82.06	31.36	70.98
22698	286.73	90.38	147.82	160.71	70.97
24053	35.17	15.83	56.32	29.72	70.97
6796	190.54	59.10	259.72	68.76	70.97
5474	783.73	210.50	556.59	211.13	70.96
22820	199.69	49.08	296.30	123.01	70.89
21796	666.75	117.72	898.08	374.38	70.88
25747	41.52	18.23	88.80	66.31	70.85
5443	12.36	13.98	34.29	27.55	70.84
12965	104.50	38.53	152.54	54.48	70.80
12332	602.96	147.89	428.95	204.34	70.77
3773	20.24	15.61	47.49	38.34	70.75
5990	310.53	65.78	375.84	76.70	70.73
18302	132.49	67.39	63.56	100.58	70.70
23964	9.67	12.94	24.20	18.71	70.68
9468	65.20	31.76	38.22	29.36	70.68
16631	14.50	12.00	57.40	80.51	70.66
21653	224.19	41.69	303.28	104.55	70.63
9097	272.42	76.73	194.27	78.59	70.61
11259	79.73	54.95	259.20	287.87	70.60
1081	515.94	100.86	394.77	126.28	70.58
18360	214.50	54.30	161.55	58.13	70.58
4789	35.91	18.80	60.76	25.85	70.56
1798	346.08	90.64	258.66	95.94	70.56
25089	69.92	34.04	119.62	57.66	70.55
24234	170.64	48.79	257.81	150.22	70.55
23270	197.31	46.26	264.06	80.46	70.54
8339	457.12	115.09	336.46	156.58	70.52
4119	104.13	29.50	142.73	41.65	70.50
18581	239.39	63.97	323.89	102.81	70.48
8188	429.99	130.68	311.19	123.41	70.46
17950	60.85	21.71	84.23	24.05	70.45
11967	1829.61	479.72	1293.20	638.27	70.45
5252	13.79	9.94	24.05	13.59	70.44
22928	235.05	79.02	162.79	64.64	70.38
16684	483.85	102.78	662.86	218.95	70.37
1463	562.00	123.95	959.70	610.46	70.36
8495	174.57	44.41	233.96	68.66	70.36
2195	94.72	37.50	60.02	37.89	70.35
3042	289.42	103.30	458.48	192.79	70.34
15330	96.55	24.69	69.31	26.59	70.32
12399	80.36	27.26	110.73	35.05	70.32
16351	67.79	22.25	109.53	54.61	70.27
3822	871.73	175.65	1184.82	469.53	70.26
21025	483.29	121.39	358.54	123.38	70.26
6548	93.61	35.75	138.74	52.86	70.23
12561	159.96	46.53	108.93	51.49	70.19
5481	36.03	51.29	102.88	72.89	70.19

TABLE 5:	GENERAL -							
		1	ı	Doc. N	<u> 0. 17988</u>			
ldendliter	MonToxMean	DEX TION	ToxiMean	ToxSD	LD/AScc			
3430	414.54	95.45	587.65	251.37	70.17			
26335	940.03	330.95	625.84	337.88	70.16			
352	77.57	33.06	132.92	75.36	70.14			
23044	213.44	34.48	253.69	53.74	70.13			
17161	1069.69	220.67	1639.92	740.76	70.11			
14352	179.82	26.18	211.25	38.48	70.10			
21993	71.93	19.33	98.79	30.29	70.09			
16756	165.09	40.66	231.09	68.53	70.09			
7537	240.15	64.50	185.74	68.65	70.09			
15986	336.87	70.00	240.73	102.05	70.07			
17256	428.03	84.39	329.02	149.16	70.04			
18151	1182.11	241.37	915.26	244.04	70.03			
18354	372.44	129.56	548.59	224.71	70.03			
19152	155.28	37.87	219.58	85.40	70.01			
8314	44.66	24.23	401.40	1027.58	70.01			
13222	132.87	25.87	162.46	38.58	69.99			
3808	157.93	29.12	224.59	103.04	69.99			
25705	432.30	81.33	560.17	181.12	69.98			
4360	341.32	51.71	279.62	90.37	69.97			
15904	48.14	15.72	70.45	28.37	69.96			
3733	307.48	109.96	502.42	242.20	69.95			
12349	248.84	51.51	206.79	57.13	69.94			
6039	293.57	52.57	404.95	147.79	69.94			
16394	529.95	207.80	998.31	642.96	69.92			
1340	194.50	26.68	172.82	51.16	69.92			
13393	68.65	39.60	120.92	67.73	69.91			
26119	115.29	35.39	165.90	63.73	69.91			
21471	-15.71	27.91	21.66	45.38	69.91			
498	513.15	100.54	694.75	220.14	69.89			
19	324.56	59.40	415.19	125.18	69.89			
22599	40.60	18.16	65.55	34.14	69.84			
7427 16520	235.31 82.42	43.69	295.43 151.39	88.56 91.91	69.83 69.83			
15642	389.37	82.84	518.22	172.58	69.83			
1430	171.08	83.75	102.96	99.73	69.82			
			61.83		00.00			
7918 13633	36.73 276.86	13.76 77.72	463.55	31.90 259.47	69.82 69.80			
7936	155.24	33.42	122.46	38.98	69.79			
15004	132.61	42.50	344.40	385.03	69.78			
15955	791.51	166.06	571.10	228.60	69.77			
1478	366.90	66.96	278.05	99.94	69.75			
7622	55.21	18.42	74.69	23.71	69.75			
22796	7.01	12.41	21.99	21.24	69.73			
17401	821.97	205.41	1398.10	790.55	69.73			
7888	262.03	59.59	349.92	94.63	69.72			
13392	181.47	36.33	245.35	76.20	69.70			
22101	199.60	77.45	118.24	102.76	69.70			
18	61.29	34.62	98.91	45.09	69.69			
	407.16	79.73	529.18	177.19	69.69			
20414	124.45	34.92	93.89	36.15	69.68			
		23.02	80.54	43.90	69.64			
	40.94	17.34	63.44	29.74	69.64			
15011		29.75	122.02	40.98	69.63			
3434		104.25	456.96	211.34	69.62			
21444		27.61	65.55	77.70	69.62			

	General		Alty. Dod	DOG. [X	921-5039VVO 0. 1793397.1
ldendifter	NonToxMean	NonToxSD	ToxiMean	ToxSD	LDAScore
3493	56.09	16.33	78.09	27.67	69.58
13727	133.95	49.99	86.22	54.51	69.58
17339	2512.73	596.97	1882.04	680.16	69.56
6518	108.28	26.66	143.55	38.12	69.55
14484	468.99	102.51	371.12	113.87	69.50
45	184.62	61.15	131.00	98.51	69.50
4235	383.35	65.78	479.39	109.87	69.47
2350	631.18	75.04	733.81	127.04	69.47
20816	359.20	71.33	589.66	361.06	69.47
20448	51.14	15.80	96.53	91.69	69.45
3608	354.31	96.17	240.45	122.08	69.45
20829	754.23	187.16	1139.64	525.74	69.43
14388	133.84	46.32	189.57	73.63	69.41
13974	269.51	60.19	455.96	331.84	69.41
13611	289.40	97.62	194.81	124.78	69.39
9452	109.85	33.54	243.69	304.19	69.39
19679	744.23	156.00	548.50	220.11	69.38
23471	80.62	27.27	134.19	61.98	69.38
15596	200.04	61.45	269.73	83.16	69.38
17159	662.06	139.21	916.50	373.05	69.37
9114	907.26	198.07	711.93	216.95	69.36
7690	188.30	76.58	286.44	101.61	69.30
4462	896.55	240.26	700.53	283.12	69.30
15146	117.66	45.74	219.43	151.19	69.28
4747	52.09	18.56	76.30	30.54	69.28
4463	171.78	48.48	116.71	64.25	69.26
21275	208.39	53.25	293.29	120.48	69.26
22537	314.16	116.85	217.64	134.70	69.24
21015	224.01	149.40	523.06	502.39	69.21
14184	104.95	35.03	152.30	77.88	69.21
16859	113.00	39.98	171.46	70.52	69.20
13359	18.79	19.20	41.55	29.30	69.17
24192	65.10	27.46	107.06	63.04	69.17
22357	537.91	143.42	683.30	219.11	69.15
22540	1928.78	500.80	1396.93	518.89	69.15
15111	1008.00	256.31	739.45	278.04	69.15
23128	629.03	117.68	522.92	126.85	69.13
9905	702.23	115.37	558.07	156.97	69.13
23387	23.75	24.01	57.01	48.92	69.12
21797	316.03	77.43	447.83	179.03	69.12
20457	401.15	78.89	312.93	100.71	69.12
13954	1425.24	337.03	1039.20	391.69	69.12
2059	134.19	33.99	173.66	52.56	69.11
21125	163.43	45.61	120.34	59.14	69.08
4048	-12.32	8.61	25.98	74.75	69.08
13349	122.00	33.87	171.03	54.95	69.08
20086	103.42	54.54	186.02	112.44 72.25	69.07
7414	177.49	39.38	246.64		69.07
4327	84.49	26.30	133.01	68.89	69.07
19011	389.45	84.42	492.01	130.94	69.04
6384	55.77	19.77	79.31	27.28	69.03
8221	192.95	91.60	123.84	72.63	69.02
11876	100.42	32.61	145.49	59.19	69.00
275	540.79	117.63	450.26	289.26	69.00
19940	19.25	13.33	32.95	17.53	68.99

	Ceneral :		Aity. Doc	ko: No. 44 Doc. N	921-5009W0 10. 1798897.1
ldendifter '	NonToxMean	NonToxSD	ToxiMean	Toxsd	LDAScore
21895	415.77	107.68	527.63	122.68	68.99
6674	2589.26	592.43	1970.28	559.33	68.97
4330	519.93	146.26	393.68	177.88	68.97
1698	59.46	33.76	154.33	156.35	68.96
6927	351.13	88.63	276.07	87.17	68.95
15879	389.05	88.44	309.88	89.07	68.94
17269	669.21	164.65	510.44	183.69	68.94
809	38.57	18.55	73.18	50.10	68.93
25567	429.75	132.83	636.76	280.62	68.91
6711	46.47	22.83	68.95	25.01	68.91
25777	370.17	112.15	585.45	351.28	68.90
22801	1309.01	233.61	1102.68	241.27	68.88
17447	912.03	226.33	708.62	297.79	68.88
1603	652.04	149.41	516.80	165.06	68.85
20460	363.41	114.33	257.61	112.54	68.84
21145	216.97	60.61	173.14	61.10	68.83
25453	225.18	45.75	181.00	63.32	68.81
14670	1156.99	219.65	1505.97	540.37	68.80
19623	56.27	29.98	87.15	39.31	68.80
12716	167.13	43.68	133.50	41.78	68.80
24236	75.95	22.54	104.28	32.01	68.79
15617	18.52	33.89	39.26	21.38	68.79
3925	498.78	107.84	390.92	113.69	68.77
20449	39.97	25.35	119.71	151.16	68.76
21390	89.77	28.23	115.10	33.69	68.76
23514	434.32	164.14	307.02	151.33	68.74
20849	259.38	58.40	370.11	145.72	68.74
794	224.09	52.71	162.86	64.32	68.71
4592	183.27	30.19	222.63	53.46	68.70
13614	325.22	70.14	423.70	138.23	68.70
12673	32.94	15.97	60.47	35.35	68.69
3125	347.43	89.73	270.34	103.82	68.67
4232	131.63	36.97	170.35	127.98	68.67
1399	187.08	41.04	289.41	156.72	68.66
13930	114.32	44.14	212.08	140.70	68.65
5689	9.09	13.75	27.68	24.99	68.63
2370	1158.46	172.59	949.24	224.94	68.63
4933	93.92	113.34	358.13	386.59	68.63
406	374.38	68.18	306.59	91.47	68.62
22957	165.92	62.59	266.91	150.55	68.62
2768	2026.25	370.26	1657.86	463.04	68.61
24197	243.13	77.12	185.53	85.38	68.60
16650	242.10	65.10	338.01	127.40	68.57
8085	36.42	17.42	58.09	26.97	68.57
1712	167.50	36.63	216.60	72.08	68.57
5565	407.17	100.35	305.41	133.21	68.56
16883	1543.18	301.38	1229.72	308.97	68.55
13622	45.42	26.78	87.87	76.11	68.55
17807	711.63	133.84	948.59	316.04	68.55
8496	39.06	20.93	64.65	32.09	68.53
11559	615.91	118.47	505.06	115.82	68.53
19094	1013.42	192.17	1274.36	327.89	68.53
7584	99.66	69.21	216.92	170.84	68.53
457	268.02	49.06	378.26	164.79	68.53
21105	323.39	54.03	270.87	64.06	68.52

18900 44 16709 81 1993 20 13348 1 11454 22 18606 66 2986 33 15644 13 17905 96 408 22 1409 22 1409 22 1583 22 1583 22 1583 22 1583 22 1596 10 1583 23 15196 10 1583 25 1583 25 1584 37 17905 36 11164 55 10887 76 15540 36	449.75 358.46 36.13 16.50 223.14 308.53 39.62 338.83 613.59 02.96 01.66 6.43 21.16 14.88 00.77 25.58 1.59 371.15 32.49 6.37 6.37	NonToxSD 84.06 172.57 15.23 39.71 54.78 116.39 12.73 225.09 853.73 243.00 67.43 35.21 195.27 78.45 27.25 12.08 14.82 66.13 169.29 129.79 24.73		130.48 176.49 37.27 84.06 133.93 267.15 18.98 345.24 879.66 232.60 78.36 45.92 258.13 93.71 47.14 25.12 17.88 105.05 199.84 153.36	6. 1798397.1 LDAScore # 68.48 68.46 68.43 68.43 68.43 68.43 68.43 68.39 68.39 68.39 68.37 68.37 68.36 68.36 68.33 68.32 68.31 68.30 68.29 68.28
18900 4 16709 8 1993 2 13348 1 11454 2 18606 6 2986 3 15644 1 22541 3 17905 9 408 2 21409 8 22543 7 3863 2 21596 1 1583 2 8917 4 17324 3 5199 6 11164 5 10887 7 15540 3	49.75 558.46 6.13 16.50 23.14 608.53 9.62 338.83 613.59 602.96 601.66 6.43 21.16 14.88 00.77 25.58 1.15 6.15	172.57 15.23 39.71 54.78 116.39 12.73 225.09 853.73 243.00 67.43 35.21 195.27 78.45 27.25 12.08 14.82 66.13 169.29 129.79 24.73	709.82 53.08 170.35 315.07 789.92 54.78 1588.45 2681.23 654.49 144.70 119.21 540.47 158.64 132.16 46.81 56.19 297.98 484.54 382.15	176.49 37.27 84.06 133.93 267.15 18.98 345.24 879.66 232.60 78.36 45.92 258.13 93.71 47.14 25.12 17.88 105.05 199.84 153.36	68.46 68.43 68.43 68.43 68.43 68.41 68.39 68.39 68.38 68.37 68.36 68.36 68.33 68.32 68.31 68.30 68.29
1993 20 13348 1 11454 20 18606 60 2986 30 15644 10 22541 30 17905 90 408 20 21409 80 22543 70 3863 20 21596 10 1583 8917 40 17324 30 5199 60 11164 50 10887 70 15540 30	26.13 16.50 223.14 508.53 39.62 338.83 6613.59 502.96 501.66 66.43 21.16 21.16 21.16 21.15 31.59 371.15 41.61 632.49 66.37	15.23 39.71 54.78 116.39 12.73 225.09 853.73 243.00 67.43 35.21 195.27 78.45 27.25 12.08 14.82 66.13 169.29 129.79 24.73	709.82 53.08 170.35 315.07 789.92 54.78 1588.45 2681.23 654.49 144.70 119.21 540.47 158.64 132.16 46.81 56.19 297.98 484.54 382.15	176.49 37.27 84.06 133.93 267.15 18.98 345.24 879.66 232.60 78.36 45.92 258.13 93.71 47.14 25.12 17.88 105.05 199.84 153.36	68.46 68.43 68.43 68.43 68.43 68.41 68.39 68.39 68.38 68.37 68.36 68.36 68.33 68.32 68.31 68.30 68.29
13348 1 11454 2: 18606 66 2986 3: 15644 1: 22541 3: 17905 9: 408 2: 21409 8: 22543 7: 3863 2: 21596 1: 1583 2: 8917 4 17324 3: 5199 6: 11164 5: 10887 7: 15540 3:	16.50 223.14 508.53 39.62 338.83 5613.59 502.96 501.66 56.43 721.16 21.4.88 00.77 25.58 11.59 571.15 541.61 532.49 66.37 66.28	39.71 54.78 116.39 12.73 225.09 853.73 243.00 67.43 35.21 195.27 78.45 27.25 12.08 14.82 66.13 169.29 129.79 24.73	170.35 315.07 789.92 54.78 1588.45 2681.23 654.49 144.70 119.21 540.47 158.64 132.16 46.81 56.19 297.98 484.54 382.15	84.06 133.93 267.15 18.98 345.24 879.66 232.60 78.36 45.92 258.13 93.71 47.14 25.12 17.88 105.05 199.84 153.36	68.43 68.43 68.43 68.43 68.41 68.39 68.39 68.37 68.37 68.36 68.36 68.33 68.32 68.31 68.30 68.29
11454 2: 18606 66 2986 3: 15644 1: 22541 3: 17905 9: 408 2: 21409 8: 22543 7: 3863 2: 21596 1: 1583 2: 8917 4 17324 3: 5199 6: 11164 5: 10887 7: 15540 3:	223.14 508.53 59.62 338.83 5613.59 502.96 501.66 56.43 721.16 21.16 21.16 21.16 21.15 31.59 371.15 32.49 33.49 36.37	54.78 116.39 12.73 225.09 853.73 243.00 67.43 35.21 195.27 78.45 27.25 12.08 14.82 66.13 169.29 129.79 24.73	315.07 789.92 54.78 1588.45 2681.23 654.49 144.70 119.21 540.47 158.64 132.16 46.81 56.19 297.98 484.54 382.15	133.93 267.15 18.98 345.24 879.66 232.60 78.36 45.92 258.13 93.71 47.14 25.12 17.88 105.05 199.84 153.36	68.43 68.43 68.41 68.39 68.39 68.38 68.37 68.37 68.36 68.36 68.33 68.32 68.31 68.30 68.29
11454 2: 18606 66 2986 3: 15644 1: 22541 3: 17905 9: 408 2: 21409 8: 22543 7: 3863 2: 21596 1: 1583 2: 8917 4 17324 3: 5199 6: 11164 5: 10887 7: 15540 3:	223.14 508.53 59.62 338.83 5613.59 502.96 501.66 56.43 721.16 21.16 21.16 21.16 21.15 31.59 371.15 32.49 33.49 36.37	54.78 116.39 12.73 225.09 853.73 243.00 67.43 35.21 195.27 78.45 27.25 12.08 14.82 66.13 169.29 129.79 24.73	315.07 789.92 54.78 1588.45 2681.23 654.49 144.70 119.21 540.47 158.64 132.16 46.81 56.19 297.98 484.54 382.15	133.93 267.15 18.98 345.24 879.66 232.60 78.36 45.92 258.13 93.71 47.14 25.12 17.88 105.05 199.84 153.36	68.43 68.43 68.41 68.39 68.39 68.38 68.37 68.37 68.36 68.36 68.33 68.32 68.31 68.30 68.29
2986 3: 15644 1: 22541 3: 17905 9: 408 2: 21409 8: 22543 7: 3863 2: 21596 1: 1583 2: 8917 4 17324 3: 5199 6: 11164 5: 10887 7: 15540 3:	338.83 338.83 3613.59 302.96 301.66 36.43 721.16 214.88 300.77 25.58 41.59 371.15 341.61 332.49 66.37	12.73 225.09 853.73 243.00 67.43 35.21 195.27 78.45 27.25 12.08 14.82 66.13 169.29 129.79 24.73	54.78 1588.45 2681.23 654.49 144.70 119.21 540.47 158.64 132.16 46.81 56.19 297.98 484.54 382.15	18.98 345.24 879.66 232.60 78.36 45.92 258.13 93.71 47.14 25.12 17.88 105.05 199.84 153.36	68.41 68.39 68.39 68.38 68.37 68.37 68.36 68.36 68.33 68.32 68.31 68.30 68.29
15644 13 22541 36 17905 96 408 26 21409 86 22543 76 3863 26 21596 16 1583 26 8917 4 17324 37 5199 66 11164 56 10887 76 15540 36	338.83 613.59 902.96 901.66 66.43 721.16 914.88 90.77 95.58 971.15 941.61 932.49 96.37	225.09 853.73 243.00 67.43 35.21 195.27 78.45 27.25 12.08 14.82 66.13 169.29 129.79 24.73	1588.45 2681.23 654.49 144.70 119.21 540.47 158.64 132.16 46.81 56.19 297.98 484.54 382.15	345.24 879.66 232.60 78.36 45.92 258.13 93.71 47.14 25.12 17.88 105.05 199.84 153.36	68.39 68.39 68.38 68.37 68.37 68.36 68.36 68.33 68.32 68.31 68.30 68.29
22541 30 17905 90 408 20 21409 80 22543 72 3863 20 1583 20 1583 20 17324 30 5199 60 11164 50 15540 30	613.59 602.96 601.66 66.43 621.16 614.88 600.77 65.58 61.59 671.15 641.61 632.49 66.28	853.73 243.00 67.43 35.21 195.27 78.45 27.25 12.08 14.82 66.13 169.29 129.79 24.73	2681.23 654.49 144.70 119.21 540.47 158.64 132.16 46.81 56.19 297.98 484.54 382.15	879.66 232.60 78.36 45.92 258.13 93.71 47.14 25.12 17.88 105.05 199.84 153.36	68.39 68.38 68.37 68.37 68.36 68.36 68.33 68.32 68.31 68.30 68.29
17905 90 408 20 21409 80 22543 70 3863 20 21596 10 1583 20 8917 4 17324 30 5199 60 11164 50 10887 70 15540 30	002.96 001.66 66.43 (21.16 014.88 00.77 05.58 1.59 071.15 641.61 032.49 (6.37 66.28	243.00 67.43 35.21 195.27 78.45 27.25 12.08 14.82 66.13 169.29 129.79 24.73	654.49 144.70 119.21 540.47 158.64 132.16 46.81 56.19 297.98 484.54 382.15	232.60 78.36 45.92 258.13 93.71 47.14 25.12 17.88 105.05 199.84 153.36	68.38 68.37 68.37 68.36 68.36 68.33 68.32 68.31 68.30 68.29
408 20 21409 80 22543 72 3863 2 21596 10 1583 20 8917 4 17324 3 5199 6 11164 5 10887 70 15540 30	201.66 66.43 721.16 214.88 00.77 25.58 1.59 671.15 641.61 632.49 76.37	67.43 35.21 195.27 78.45 27.25 12.08 14.82 66.13 169.29 129.79 24.73	144.70 119.21 540.47 158.64 132.16 46.81 56.19 297.98 484.54 382.15	78.36 45.92 258.13 93.71 47.14 25.12 17.88 105.05 199.84 153.36	68.37 68.37 68.36 68.36 68.33 68.32 68.31 68.30 68.29
21409 80 22543 73 3863 2 21596 10 1583 25 8917 4 17324 3 5199 66 11164 55 10887 70 15540 30	66.43 (21.16 (21.4.88 (00.77 (5.58 (1.59 (71.15 (41.61 (32.49 (6.37	35.21 195.27 78.45 27.25 12.08 14.82 66.13 169.29 129.79 24.73	119.21 540.47 158.64 132.16 46.81 56.19 297.98 484.54 382.15	45.92 258.13 93.71 47.14 25.12 17.88 105.05 199.84 153.36	68.37 68.36 68.36 68.33 68.32 68.31 68.30 68.29
22543 7: 3863 2: 21596 10 1583 2: 8917 4 17324 3: 5199 66 11164 5: 10887 70 15540 3:	721.16 14.88 00.77 25.58 1.59 371.15 341.61 332.49 6.37	195.27 78.45 27.25 12.08 14.82 66.13 169.29 129.79 24.73	540.47 158.64 132.16 46.81 56.19 297.98 484.54 382.15	258.13 93.71 47.14 25.12 17.88 105.05 199.84 153.36	68.36 68.36 68.33 68.32 68.31 68.30 68.29
3863 2 21596 10 1583 20 8917 4 17324 3 5199 6 11164 5 10887 70 15540 30	214.88 00.77 25.58 1.59 371.15 341.61 32.49 6.37	78.45 27.25 12.08 14.82 66.13 169.29 129.79 24.73	158.64 132.16 46.81 56.19 297.98 484.54 382.15	93.71 47.14 25.12 17.88 105.05 199.84 153.36	68.36 68.33 68.32 68.31 68.30 68.29
21596 10 1583 25 8917 4 17324 3 5199 66 11164 55 10887 70 15540 36	00.77 25.58 11.59 371.15 341.61 32.49 6.37	27.25 12.08 14.82 66.13 169.29 129.79 24.73	132.16 46.81 56.19 297.98 484.54 382.15	47.14 25.12 17.88 105.05 199.84 153.36	68.33 68.32 68.31 68.30 68.29
1583 2: 8917 4: 17324 3: 5199 6: 11164 5: 10887 7: 15540 3:	25.58 11.59 171.15 141.61 132.49 16.37	12.08 14.82 66.13 169.29 129.79 24.73	46.81 56.19 297.98 484.54 382.15	25.12 17.88 105.05 199.84 153.36	68.32 68.31 68.30 68.29
8917 4 17324 3 5199 6 11164 5 10887 7 15540 3	1.59 671.15 641.61 632.49 6.37	14.82 66.13 169.29 129.79 24.73	56.19 297.98 484.54 382.15	17.88 105.05 199.84 153.36	68.32 68.31 68.30 68.29
17324 3 5199 6 11164 5 10887 7 15540 3	671.15 641.61 632.49 6.37 66.28	66.13 169.29 129.79 24.73	297.98 484.54 382.15	105.05 199.84 153.36	68.30 68.29
17324 3 5199 6 11164 5 10887 7 15540 3	641.61 632.49 6.37 6.28	169.29 129.79 24.73	484.54 382.15	105.05 199.84 153.36	68.30 68.29
5199 6- 11164 5: 10887 7: 15540 3:	641.61 632.49 6.37 6.28	169.29 129.79 24.73	484.54 382.15	199.84 153.36	68.29
11164 53 10887 70 15540 30	32.49 6.37 6.28	129.79 24.73	382.15	153.36	
10887 70 15540 30	6.37 6.28	24.73			IOO.ZO
15540 30	6.28		104.29	25.72	68.28
		11.51	64.56	42.69	68.27
	162.93	273.54	892.52	329.00	68.26
	96.29	101.41	489.27	121.68	68.26
		20.84	101.53	44.09	68.25
		63.33	465.71	138.80	68.25
		27.49	166.56	218.99	68.25
		53.69	235.92	70.52	68.24
		113.88	332.58	104.75	68.23
	31.45	106.61	540.94	147.67	68.23
		48.35	197.32	125.83	68.22
	02.48	30.70	135.61	50.44	68.21
		22.52	56.30	38.43	68.21
		42.28	60.83	59.54	68.21
		31.11	94.44	49.63	68.20
		20.73	98.29	25.44	68.20
	0.66	16.63	56.30	35.67	68.19
			631.19		68.19
		79.30	558.85	188.65	68.18
		55.30	189.70	64.51	68.17
		31.03	47.27	32.31	68.17
		32.97	104.50	38.05	68.15
		45.05	251.81	139.06	68.15
		18.28	64.87	29.10	68.14
		27.42	52.36	35.41	68.14
		40.29	131.23	57.33	68.11
		196.05	591.05	213.64	68.11
		9.54	24.57	23.36	68.10
		18.24	47.84	25.93	68.10
			1569.20	25.93 595.26	
		515.11			68.10
		19.78	90.90	31.53	68.09
		47.01	302.78	72.37	68.09
		191.34	549.12	221.95	68.09
		159.14 47.15	630.49 189.99	196.22 56.21	68.09 68.08

TABLE 5:	<u>Ceneral</u>		Ality. Docket No. 44921-50391			
0.0. 2020				1 1-742 1 1 1 1 1 1 1 1 1 1 1 1	lo. 179839	
ldentifier	NonToxMean		ToxiMean	ToxSD	LDASco	
1942	12.95	13.20	45.54	48.73	68.08	
514	4.47	50.55	52.66	58.22	68.08	
19768	683.61	138.87	883.43	263.72	68.06	
5183	204.36	51.85	284.20	109.64	68.06	
24375	107.65	26.73	157.63	66.73	68.05	
6059	199.74	47.55	169.09	54.88	68.04	
12937	20.23	21.04	62.89	58.77	68.04	
3245	97.45	32.09	132.43	48.29	68.02	
19469	376.00	72.62	300.86	98.78	68.02	
22696	72.56	48.08	25.09	39.70	68.02	
4355	116.49	44.14	163.32	82.62	68.01	
21579	110.85	35.32	153.51	68.22	68.00	
1431	521.93	166.42	374.89	194.57	67.99	
9673	66.91	27.02	44.90	26.55	67.99	
20257	137.10	42.30	102.53	48.53	67.99	
12961	185.53	42.22	151.73	41.05	67.97	
22538	338.12	80.15	255.39	90.87	67.97	
7243	56.76	22.51	79.45	28.02	67.96	
5634	64.78	29.00	96.85	47.59	67.96	
17438	62.59	33.01	31.95	40.59	67.96	
1581	70.58	17.49	93.61	30.37	67.96	
25379	75.47	18.52	100.15	31.59	67.95	
2153	153.35	49.73	361.58	346.41	67.94	
7499	17.61	8.98	28.61	16.07	67.94	
6102	150.29	40.49	187.75	46.70	67.94	
3878	479.08	106.98	389.02	107.79	67.93	
20122	174.65	39.83	214.44	52.37	67.93	
6828	122.12	53.45	207.48	104.75	67.92	
11455	106.22	31.64	162.96	86.72	67.92	
16058	146.84	40.61	225.40	102.56	67.92	
23567	42.81	36.02	114.42	124.08	67.90	
19998	290.74	86.07	214.07	102.16	67.90	
24054	25.17	14.49	39.46	20.61	67.89	
22352	139.84	54.27	253.98	211.46	67.89	
21146	119.46	36.34	92.66	39.95	67.89	
11791	136.74	46.57	173.47	50.00	67.88	
19086	102.38	32.93	153.92	65.34	67.87	
13111	235.97	68.33	187.10	77.46	67.86	
14959	595.43	107.90	761.75	241.85	67.86	
22103	222.38	50.69	294.15	83.02	67.84	
1454	110.82	42.51	160.57	70.87	67.83	
1844	167.38	35.91	216.33	60.90	67.83	
13023	110.03	149.11	607.48	946.55	67.81	
22487	34.23	15.90	60.56	35.71	67.80	
7543	268.34	96.90	185.92	86.13	67.80	
3107	779.30	164.38	696.29	236.36	67.80	
8975	102.89	42.50	68.46	48.06	67.80	
825	59.44	21.82	42.35	25.50	67.80	
4473	192.83	43.69	148.92	66.25	67.78	
18109	15.28	14.01	39.23	36.39	67.78	
		103.22			67.76	
2845 11974	651.40 247.17	76.60	768.87 176.82	160.95 86.10	67.73	
3547	14.36	11.57	25.42	14.68	67.73	
22931 7161	87.38 46.95	47.83 14.21	52.96 64.96	42.09 24.05	67.72 67.69	

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TABLE 5: CENERÁL AGy. Docket No. 44921-5089V Doc. No. 179889							
ldentifier	MonToxMean	Dexotroly	ToxXVean	ToxSD	LDAScore		
9061	1135.34	236.25	942.16	326.87	67.69		
12331	618.37	134.26	483.61	182.86	67.69		
13962	523.52	110.90	438.13	148.62	67.68		
24277	134.73	41.18	181.65	55.29	67.67		
14790	176.36	78.47	102.86	72.99	67.67		
18528	361.23	136.11	589.25	331.81	67.66		
19665	81.08	33.19	142.12	76.68	67.66		
14242	32.67	14.37	49.07	20.85	67.64		
17407	1713.79	374.66	1443.40	337.67	67.64		
6765	820.06	183.88	653.70	231.97	67.62		
9514	675.01	129.64	570.37	132.48	67.62		
22602	334.13	94.17	237.80	117.01	67.57		
19822	1669.92	376.25	1281.37	430.75	67.56		
9699	58.47	18.16	42.68	19.82	67.55		
12812	99.34	33.83	74.64	38.83	67.55		
24566	200.15	82.45	135.98	77.12	67.54		
17499	24.56	23.09	55.11	35.51	67.53		
18447	1233.88	274.38	958.23	329.75	67.53		
21014	142.85	40.94	210.35	117.97	67.52		
2536	406.73	114.26	323.04	150.73	67.51		

TABLE 5A:	ACYCLOMR s): 24, 163 hrs			DOC.	4921-5039W0 No. 1793397.1
ldendifer	MonToxMean	NonToxSD	ToxMean	Toxsd	LDAScore
25419	159.17	64.32	38.16	8.66	99.48
25415	169.94	67.00	36.68	10.41	99.48
1872	723.76	219.40	182.16	67.65	99.36
17198	814.88	264.22	145.60	82.51	99.14
17567	1172.88	345.04	2623.71	250.73	98.97
14959	616.48	139.00	1288.62	137.24	98.84
20945	896.18	242.10	1896.41	180.47	98.80
16245	387.31	114.26	41.23	45.70	98.80
10887	73.83	25.90	17.83	4.94	98.67
4222	544.00	97.71	1042.92	176.22	98.67
19161	1064.06	296.13	2360.57	327.19	98.63
15626	1281.17	347.65	3250.17	648.92	98.58
11849	701.03	165.94	1608.51	320.36	98.50
20872	832.75	216.84	1896.38	376.01	98.50
15875	1175.87	375.74	2630.93	347.24	98.45
10498	1085.24	333.50	2568.66	334.73	98.45
20884	639.33	257.31	46.55	54.58	98.37
13151	702.17	319.88	2212.94	511.24	98.32
18611	1409.99	440.83	2945.57	350.22	98.28
20885	538.89	188.67	83.65	54.88	98.28
16244	40.33	21.14	-0.26	5.01	98.24
16918	1172.71	382.43	2977.59	742.16	98.15
3027	1121.82	319.93	2267.04	262.71	98.15
20056	309.42	59.71	118.77	41.29	98.11
16205	932.31	252.05	1896.76	307.48	98.02
20812	1107.00	305.73	2296.29	293.56	97.98
14384	375.87	76.79	604.80	58.76	97.94
24615	809.56	214.82	1799.64	368.34	97.94
17524	1175.90	282.46	541.39	119.09	97.85
20839	1037.63	274.92	2191.65	391.42	97.85
18250	1088.85	307.06	2058.50	257.28	97.77
22846	1492.71	284.92	855.87	120.77	97.68
3026	512:85	99.35	933.31	159.19	97.64
13647	864.69	256.24	2095.09	431.00	97.64
17563	1195.18	331.95	2542.06	387.04	97.59
17473	422.91	93.66	796.08	169.41	97.59
20746	673.25	126.38	1181.14	211.43	97.59
19359	1061.61	314.08	2336.47	601.88	97.51
15201	1478.32	513.09	3558.35	808.75	97.47
15052	1433.93	492.46	3597.61	980.31	97.42
4490	76.73	73.42	270.49	135.30	97.42
2696	773.00	225.06	1860.28	390.37	97.42
22552	313.47	90.91	645.46	155.87	97.38
1694	1139.80	326.74	2255.77	345.62	97.38
19824	224.99	67.29	73.65	27.54	97.34
16333	133.41	38.84	45.56	15.40	97.16
16150	495.73	119.09	217.15	50.78	97.16
15928	157.91	59.34	354.44	52.65	97.16
15335	424.77	103.56	772.54	127.56	96.99
15202	788.60	313.98	1866.58	266.92	96.99
18749	116.41	42.26	34.59	11.87	96.95
16164	1075.08	276.06	1925.19	293.15	96.86
20088	384.13	78.71	185.57	40.67	96.78
23989	1057.77	294.07	466.66	116.46	96.74
24048	695.95	214.73	269.65	68.66	96.65
4254	144.21	377.79	15.39	14.62	96.65

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Timepoint	: ACYCLOVIR (s)): 24, 163 hrs			<u>Doc.</u>	1921-5039VVO No. 1793397.1
ldendifer	NonToxMean	NonToxSD -	ToxiMean :	TOXSD	LDAScore
24577	1170.09	364.08	2400.41	380.56	96.65

TABLE SE	: ACYCLOVIR (s): 6 lins		Ally, Do		44921-5039000		
					o. 1793397.1		
ldendifier					LDAScore:		
2572	1079.22	272.72	292.63	34.53	99.96		
657	325.86	85.70	825.82	55.34	99.96		
15174	582.65	154.01	1160.30	40.29	99.96		
22060	120.39	46.22	489.85	39.32	99.87		
21151	108.89	64.20	1068.35	133.62	99.87		
17546	542.45	131.44	1300.18	82.92	99.83		
8477	528.52	158.91	1301.77	46.70	99.83		
16993	147.03	72.47	5.04	12.16	99.79		
10667	41.39	38.71	-236.50	27.34	99.79		
3822	915.99	266.90	2473.92	69.91	99.74		
17157	50.71	97.99	182.37	23.19	99.70		
4532	266.41	75.57	62.22	7.77	99.70		
15004	160.24	140.85	1576.85	132.63	99.70		
4832	149.39	47.92	32.65	11.70	99.70		
11836	219.06	64.52	69.22	8.46	99.61		
24390	165.64	111.19	-250.24	60.63	99.61		
24200	421.04	138.48	1131.90	38.93	99.61		
15002	137.74	85.32	904.65	69.55	99.61		
3713	1141.71	271.69	457.32	36.48	99.57		
6778	109.70	31.47	262.53	25.12	99.57		
7936	149.06	36.24	49.64	5.41	99.57		
13542	446.95	133.65	59.60	28.78	99.57		
22385	107.82	53.29	22.50	3.64	99.57		
20700	74.63	363.62	3153.73	549.12	99.57		
15190	1801.09	1118.86	7715.59	404.27	99.53		
20698	-0.10	63.70	407.01	114.99	99.53		
9757	408.35	98.75	159.69	13.35	99.53		
15077	87.92	41.66	6.25	3.82	99.53		
12301	58.09	29.22	219.75	30.79	99.53		
3304	881.73	218.32	247.94	48.65	99.53		
24041	11.98	28.98	230.57	61.01	99.53		
19780	47.97	40.89	251.97	31.79	99.53		
12899	-9.55	17.51	146.15	30.06	99.49		
23387	29.40	30.63	203.21	32.17	99.49		
20699	86.60	192.11	2061.40	288.42	99.49		
606	-48.26	30.91	77.22	17.44	99.49		
8874	118.63	59.95	395.15	33.95	99.49		
5475	445.78	146.14	115.95	22.16	99.49		
985	21.00	43.98	281.44		99.49		
24243	258.96	69.54	71.35	13.38	99.44		
6057	104.47	44.96	358.76	52.20	99.44		
1892	-6.30	35.30	58.25	24.52	99.40		
275	519.62	162.35	80.95	30.90	99.40		
2695	210.32	124.89	613.77	27.62	99.40		
7804	1764.46	393.34	898.61	69.66	99.40		
20701	32.96	54.09	404.95		99.40		
12420	23.45	48.04	-131.74		99.36		
15003	33.26	83.63	776.16		99.36		
24564	568.95	174.41	66.56	19.02	99.36		
24246	300.44	96.46	678.79	40.80	99.36		
9423	939.03	252.01	2035.02	137.31	99.36		
23151	380.64	125.08	82.73		99.36		
9071	48.03	25.31	2.95	1.90	99.36		
2905	244.95	103.58	780.68	56.20	99.36		

TABLE 58	: ACYCLOVIR		Ally, Dog	ket No. 449	921-5089VVO			
	Timepoint(s): 6 hrs Doc. No. 1793397.							
ldendifier	MonToxMean	NonTox&D	ToxMean	ToxSD	LDAScore			
5931	182.37	66.37	9.61	8.36	99.36			
24042	18.74	67.93	772.42	198.67	99.36			
17765	1285.78	452.80	2557.67	63.52	99.31			
17470	102.21	54.04	410.26	61.20	99.31			
2010	32.91	314.74	355.60	90.97	99.31			
20848	510.27	150.82	1095.65	24.39	99.31			
17591	385.36	90.04	744.25	62.21	99.27			
1340	192.09	49.88	96.02	6.73	99.27			
7806	51.84	19.09	122.32	8.65	99.27			
7493	78.12	36.73	174.58	4.92	99.27			
20035	180.04	101.54	817.96	80.03	99.23			
19657	3.30	17.67	125.30	19.29	99.23			
24563	309.99	99.44	4.82	17.01	99.23			
22453	171.10	46.39	60.55	10.10	99.23			
23995	161.66	57.89	380.58	31.88	99.23			
15191	1989.62	1126.31	8988.79	1157.81	99.23			
11326	332.40	95.69	94.87	14.33	99.23			
7586	793.77	201.98	306.02	29.21	99.23			
2392	154.00	87.41	1138.50	400.02	99.23			
4205	222.66	72.63	591.81	54.59	99.23			
8245	54.69	20.70	113.91	5.63	99.23			
16324	194.65	61.24	42.04	11.21	99.23			
12404	105.63	62.52	662.04	167.20	99.23			
7639	753.14	166.01	339.14	44.82	99.23			
20895	331.67	100.56	50.78	24.52	99.19			
22018	158.66	41.98	360.67	50.74	99.19			
3823	524.10	147.80	1250.81	97.60	99.14			
6477	13.58	161.23	99.71	26.48	99.14			
223	11.18	17.65	124.61	25.10	99.14			
15146	130.31	69.77	744.71	139.52	99.14			
25069	134.29	60.66	25.21	12.36	99.14			
3431	1503.29	617.93	3867.63	202.46	99.14			
3271	576.75	119.79	239.69	44.71	99.14			
6054	26.29	47.50	386.77	134.53	99.14			
20202	627.69	194.14	49.29	62.88	99.14			
25546	473.74	139.13	145.65	40.29	99.14			
19781	113.50	55.43	454.62	115.29	99.14			
22488	58.52	53.30		50.62	99.10			
10281	168.73	157.24		591.27	99.10			
9452	125.58	75.70	1560.61	720.48	99.10			
2153	182.83	143.86	1201.51	175.46	99.10			
13745	27.62	27.50	297.57	96.36	99.10			
1218	198.28	37.55	92.30	12.70	99.10			
15042	63.76	42.44	353.57	60.19	99.10			
9521	95.05	24.96	222.10	33.80	99.10			
90Z I	30.00	24.90	222.10	აა.o∪	J3. IU			

TABLE 5C Timepoint	: ADR (S): 120, 163 hrs		Ally. Do	eket No. 44 Doc. 1	1921-5039WO lo. 1793397.1
	MonToxMean		ToxMean		
1688	5353.71	3830.70	26.75	39.23	99.44
25469	1577.10	747.53	25.91	13.98	99.44
1684	2831.22	1612.49	24.53	31.27	99.44
17829	2235.92	1102.19	68.14	58.70	99.44
25468	2186.32	1123.33	11.47	13.96	99.44
26150	783.82	532.35	-38.06	15.15	99.40
1687	1894.26	855.11	45.95	32.94	99.31
1685	9226.22	7079.05	58.61	94.83	99.31
1689	4411.94	2221.14	43.26	28.59	99.31
17832	1976.64	870.20	17.22	14.67	99.18
19358	792.56	328.04	6.04	46.93	98.71
18907	102.84	49.44	17.51	7.82	97.33
25467	636.86	169.13	1263.84	257.87	97.25
4011	457.30	195.33	953.88	244.47	96.78
14199	71.52	35.30	26.74	5.48	96.09
2852	49.29	27.79	19.72	2.87	95.92
21140	109.12	47.83	38.53	9.71	95.83
4594	63.32	30.73	22.33	5.88	95.44
7089	102.25	41.59	48.02	7.26	95.40
2984	582.52	185.78	258.99	70.15	95.27
1831	49.81	23.36	10.63	6.34	94.67
25705	455.64	115.95	612.52	59.36	94.45
16109	414.49	75.48	556.38	45.14	94.28
11165	529.31	155.75	250.39	51.44	93.98
19237	77.59	39.42	10.97	14.64	93.68
16401	1239.09	805.30	2322.72	497.80	93.68
109	547.11	293.37	1681.21	571.81	93.63
4312	77.34	39.04	127.17	10.27	93.51
16400	580.43	474.95	1090.34	298.55	93.42
18794	138.50	77.18	48.95	11.54	93.38
7489	81.80	32.11	25.62	12.02	93.34
2586	52.04	30.18	14.65	5.97	93.04
17742	1059.53	304.09	1669.92	253.17	92.99
956	54.64	39.22	1.90	10.80	92.78
17563	1201.38	349.79	1590.38	101.68	92.69
2125	78.10	82.13	-4.59	19.58	92.69
24844	32.09	22.37	8.57	2.19	92.65
16676	38.73	27.29	17.84	2.42	92.61
19189	60.60	62.41	14.86	7.74	92.61
16521	285.39	89.91	428.27	70.75	92.56
14430	34.74	32.18	3.43	3.27	92.48
1301	325.54	286.68	14.24	17.87	92.43
21006	55.98	36.26	16.33	6.58	92.39
23778	69.71	34.49	24.74	8.89	92.26
2812	193.71	46.19	284.99	38.81	92.22
16407	459.56	105.59	825.37	208.72	92.12
16426	280.85	109.78	118.35	49.48	91.79
24232	160.98	60.26	72.06	20.28	91.66
6059	193.61	50.50	116.52	21.50	91.40
17234	634.83	173.83	432.41	36.23	91.27
13025	335.88	84.38	224.82	24.51	91.23
20757	411.95	220.62	771.74	157.51	91.09
10076	105.84	50.96	140.17	10.04	91.06
4010	961.35	403.18	2024.17	546.67	91.05
	147.73	137.94	463.05	162.91	90.96
22220	1147.73				

			000 75 3	0 00 00	
	ADR		- Wity. Dog	3.391 NO. 44	1921=30:12/\/\ 10. 1798397.1
	(s)): 120, 163 hrs				Taractura de la constanta de l
	NonToxMean .		ACTION ACCOUNTS AND ACCOUNTS AN		LDAScore -
8597	240.84	58.03	315.06	30.21	90.89
16582	56.67	19.30	29.31	5.28	90.89
1321	510.50	316.91	1496.60	371.84	90.88
14337	211.54	44.52	148.64	18.05	90.84
19191	1011.62	281.12	631.62	94.67	90.76
20716	652.51	142.58	1029.68	128.61	90.75
8017	188.58	76.89	406.11	152.05	90.66
18502	670.14	243.60	313.44	111.24	90.63
2782	214.23	97.42	522.66	132.70	90.62
13354	88.85	57.76	34.85	9.49	90.50
22696	62.39	50.49	-5.66	12.94	90.50
4242	749.55	193.32	1337.17	268.80	90.49
12660	71.28	22.64	44.42	5.74	90.46
15892	18.01	22.79	36.39	7.65	90.46
25517	39.47	33.13	2.75	7.99	90.41
22697	59.68	40.43	15.77	10.73	90.33
16448	129.58	38.04	43.90	19.88	90.32
1058	75.84	35.86	30.65	8.83	90.24
7863	1200.90	307.30	1386.51	55.05	90.24
11967	1725.17	550.50	1088.97	237.35	90.20
516	55.59	33.01	17.20	9.40	90.11
25736	24.77	16.54	7.84	4.16	89.98
6544	53.15	110.52	379.03	173.46	89.89
21651	32.42	26.38	7.38	3.39	89.85
16581	39.86	19.57	17.01	3.68	89.72
2607	218.90	50.79	175.06	11.54	89.55
15247	637.26	192.10	1113.60	210.08	89.41
6691	119.56	54.62	32.51	16.76	89.41
20702	300.67	70.50	198.62	33.60	89.38
10109	1044.97	289.50	1362.38	117.72	89.34
24040	557.97	233.98	196.88	87.97	89.29
1169	90.18	28.72	92.58	4.98	89.21
5421	279.85	96.20	162.53	31.31	89.21
19244	1224.80	382.60	1678.75	173.86	89.17
3015	2307.62	994.19	2930.38	184.09	89.12
21695	60.82	36.14	4.02	14.39	89.03
11218	102.70	37.49	47.65	19.31	88.95
21766	286.32	76.11	204.08	26.91	88.95
494	698.84	215.82	1186.69	207.80	88.86
24528	61.08	25.93	19.77	8.13	88.81

PCT/US02/16173

		-387	-		
TARLE SD	AY			eket No. 4	1021-5030
	360 hrs				Vo. 17983
	NonToxMean	are the second of the second o	 Times/Moora		
6360	7.76	5.80	30.12	3.08	99.36
18826	1069.51	321.45	473.21	31.01	99.32
24886	1264.06	371.77	2165.64	63.29	99.27
6517	233.00	158.96	437.46	12.19	99.19
16576	111.85	41.47	67.47	0.61	99.19
22846	1490.00	287.46	852.52	44.29	99.10
2708	383.37	87.00	425.82	1.07	98.97
14349	413.99	144.50	244.25	2.46	98.97
13023	187.03	420.15	-22.68	5.86	98.85
6585	653.17	368.20	239.73	12.54	98.76
15093	35.54	16.49	-7.39	3.34	98.76
25066	136.51	58.55	34.09	4.82	98.72
21796 2010	696.51	210.24	1145.40	47.75	98.67
3610 24226	1195.23	334.91	780.10	28.06	98.67
24236	80.80	27.74	39.65	1.30	98.67
16156 17672	858.19	319.87	3220.95	1137.23	98.59
17672 472	2630.76	734.18	3325.88	12.22	98.59
472 15462	658.60	175.52 35.25	1275.16	135.82	98.59 98.55
	86.01 7.63		28.41	2.70	
10159 457		18.34	30.64	0.79	98.55
22093	284.29 -21.77	87.14 38.66	421.47	22.09	98.42
22093 5212		67.72	56.32	5.98	98.37
	189.25		0.93	13.47	98.33
11368 22731	20.04 24.04	24.02 24.97	29.17	0.30	98.29
23313	404.59	87.09	9.37	0.65	98.29 98.29
25178	4.72		30.15	9.22	98.25
25176 11561	l · · · · -	6.91			
14120	389.44 740.64	87.17 221.38	225.68 1485.37	12.21	98.25
4622	608.46	110.69	956.12	218.95 79.60	98.20 98.20
12894	141.35	34.03	72.48	5.99	98.12
3447	33.98	17.04	11.74	0.97	98.03
1410	99.49	46.40	23.57	5.02	97.99
22103	235.14	64.79	449.04	43.64	97.99
14185	204.58	95.56	321.83	10.61	97.99
24770	233.44	86.68	44.57	16.38	97.95
21044	24.56	22.71	-11.37	2.16	97.95
24721	203.93	53.62	144.06	2.18	97.95
20184	11.29	9.18	20.17	0.35	97.95
6062	182.74	59.18	106.49	1.72	97.90
16172	467.00	120.07	308.58	10.28	97.86
19731	225.58	214.69	63.47	15.21	97.82
16155	1075.32	410.99	3024.24	951.97	97.82
5425	85.43	33.27	153.80	5.81	97.82
1463	620.25	296.25	1592.90	460.18	97.78
2696	777.90	238.27	1766.44	260.04	97.78
2126	223.21	75.66	478.73	56.24	97.78
16245	384.42	117.13	589.67	20.00	97.73
4683	211.56	40.44	323.17	13.38	97.73
11590	276.38	78.94	238.32	1.35	97.73
556	82.41	35.72	33.35	5.40	97.69
10241	67.50	27.73	4.65	11.47	97.69
275	516.42	162.40	1038.56	193.69	97.65
10784	24.89	30.12	-21.79	6.29	97.65
6523	30.10	14.87	8.53	1.41	97.60
4467	656.82	156.93	1236.28	198.60	97.60

					KAT TO BE AN AND THE STATE OF T
	e ay				4921 -5 039/WO
	(s): 360 hrs 👢				No. 1798397.1
ldendifier	NonToxMean:	NonToxSD	Lox(Meau)	ToxSD	
24042	21.86	84.61	58.64	10.50	97.56
15135	706.76	182.37	1034.39	31.17	97.52
6790	63.12	24.62	19.35	3.71	97.52
6743	1405.82	280.92	2262.37	206.08	97.52
6351	45.29	26.36	88.65	2.94	97.48
18942	-11.45	10.37	24.19	16.98	97.43
6726	393.51	104.57	242.70	10.50	97.43
9808	30.77	14.74	50.69	1.63	97.39
21078	567.53	143.04	363.30	11.51	97.39
22619	371.47	90.93	316.17	2.78	97.39
1058	75.62	35.90	24.89	3.27	97.39
22692	209.08	63.65	106.95	5.50	97.39
21914	404.11	71.79	635.41	66.12	97.39
22063	68.58	25.03	47.50	0.52	97.35
1162	10.60	37.16	138.81	40.04	97.35
15224	689.92	152.18	1138.23	103.80	97.35
24178	2.75	9.42	20.42	4.02	97.31
20891	140.77	49.29	78.20	3.12	97.31
17393	145.61	59.87	87.56	1.63	97.31
17061	557.85	95.17	904.28	161.35	97.22
21656	54.49	25.08	22.60	1.43	97.18
23651	656.34	623.55	2317.09	936.44	97.18
2357	76.01	23.05	69.48	0.72	97.13
18130	378.79	74.65	591.90	50.88	97.13
23898	10.17	6.54	30.37	7.27	97.13
9363	89.83	29.45	164.93	13.77	97.09
2920	78.42	29.88	124.28	3.10	97.09
21930	597.55	127.69	1108.70	256.19	97.05
12770	600.59	229.99	230.85	29.99	97.05
23799	146.92	43.68	75.83	5.19	97.05
1757	23.41	13.46	-0.34	3.13	97.05
18419	1317.37	373.39	2669.33	536.46	97.05
7023	364.44	77.23	413.76	3.42	97.01
5811	35.76	18.86	52.44	0.86	97.01
1529	290.79	62.32	164.72	18.30	97.01
15259	227.54	56.59	163.79	2.45	97.01
14095	379.89	98.99	246.75	8.11	97.01
17107	2200.10	722.73	4177.24	487.81	96.96
13393	76.83	46.83	166.60	16.98	96.96
15411	305.14	91.59	162.75	9.40	96.96

	: BEA (s): 6, 24 hrs		Ally. Docket No. 44921-5039 Doc. No. 179339			
	NonToxMean		ToxMean		LDAScore.	
21011	154.85	315.31	689.68	175.67	98.32	
21015	271.80	338.52	933.97	202.57	97.98	
21013	214.58	340.35	710.95	153.21	97.68	
22057	349.60	55.08	477.85	22.42	97.33	
8477	529.15	164.21	876.56	49.54	97.03	
23849	285.73	136.63	521.70	61.30	96.65	
15969	339.58	71.02	574.34	94.47	96.47	
5901	92.67	50.17	176.54	31.37	96.43	
17034	885.84	144.51	1193.98	93.26	95.36	
23140	172.19	53.17	288.43	107.64	94.67	
6143	560.24	135.28	861.03	122.25	94.20	
22931	79.76	48.36	19.97	4.03	94.15	
13608	21.70	23.78	64.61	13.40	94.07	
3167	308.80	70.69	438.24	41.83	94.07	
1777 1	769.01	307.47	1086.30	71.95	93.98	
10477	87.73	37.06	137.33	14.74	193.42	
17563	1206.59	350.63	917.80	32.07	92.99	
3551	440.44	98.71	307.06	31.45	92.69	
22885	1347.43	493.33	2161.55	380.96	92.61	
8515	262.49	109.22	415.39	29.53	92.52	
20745	471.03	70.08	354.88	30.48	92.30	
4748	110.75	127.98	202.78	32.60	92.18	
8639	368.78	99.86	530.10	59.33	91.92	
14874	50.39	17.11	78.62	12.99	91.66	
21625	2492.78	1106.79	1806.41	173.96	91.53	
2729	638.00	170.57	841.33	45.73	91.53	
6844	124.24	58.33	32.11	26.05	91.53	
19993	2317.82	568.48	3182.90	249.42	91.36	
10742	62.40	27.77	27.67	6.47	91.32	
8205	398.00	113.24	583.05	83.76	91.23	
17400	121.53	78.34	254.55	53.39	91.23	
1698	73.83	75.94	197.90	43.39	91.22	
3557	107.65	39.24	177.58	40.40	91.14	
18905	1365.13	302.42	1751.48	71.12	91.06	
7276	72.89	27.71	30.30	10.38	91.00	
15111	954.53	281.39	581.40	91.29	90.97	
14929	800.42	430.77	1096.37	215.26	90.80	
18077	2620.17	1190.40	1912.67	180.40	90.28	
14862	181.12	72.28	270.61	28.36	90.28	
10636	369.25	95.44	241.19	39.11	90.24	
2057	291.01	74.63	398.13	39.83	90.15	
21014	155.16	80.93	268.20	58.44	90.15	
4232	137.37	62.35	64.91	22.20	90.15	
5687	189.73	55.56	278.09	42.24	90.07	
13614	342.46	95.82	447.07	32.82	90.03	
	98.56	37.96	208.10	79.38	89.97	
	2202.25	730.97	2581.41	115.51	89.72	
	29.03	14.66	50.58	9.56	89.68	
	463.39	111.03	329.66	41.65	89.68	
	29.07	15.66	43.28	5.07	89.64	
	119.24	35.99	215.47	51.98	89.63	
	38.97	27.26	103.83	29.32	89.63	
	4123.05	2816.44	2751.13	405.49	89.60	
	74,49	49.29	192.85	48.58	89.59	
	1 マ.マン	マン・ムン	1134.00	70.00	UJ.JJ	
	728.25	203.79	1236.56	236.71	89.59	

TABLE SE	: BEA	15. 60 3 41.24	AMW. Dood	kent No. 44	1921-5089WO
Vimepoint	((s)): 6, 24 hrs			Doc. R	lo. 1793397.1
ldendifer	NonToxMean 3	NonToxSD	ToxMean	ToxSD	LDAScore :
3081	387.79	84.55	253.14	37.34	89.46
7895	997.72	308.71	1503.83	263.73	89.46
3924	183.13	68.64	101.62	16.04	89.34
24181	85.08	27.22	145.70	16.84	89.33
2752	405.86	139.92	593.99	60.88	89.33
1159	886.55	231.02	611.65	64.43	89.29
24388	188.48	72.47	251.08	30.81	89.29
3926	176.00	56.73	103.17	18.22	89.25
18981	231.55	65.64	189.11	9.52	89.21
7838	19.58	15.00	21.80	3.33	89.17
24537	528.52	104.62	396.25	38.65	89.08
2688	173.21	54.36	243.95	28.16	89.08
19484	184.45	70.04	320.46	37.72	89.03
22855	561.72	144.30	886.81	141.20	89.03
12979	391.37	216.96	627.81	88.47	88.94
7223	88.64	27.62	152.72	24.63	88.94
23159	416.48	85.49	537.44	47.74	88.91
13563	1028.27	255.15	1603.62	170.06	88.86
3696	42.57	32.72	106.32	10.80	88.86
2855	920.64	212.77	1412.67	261.05	88.77
3580	9.48	10.46	22.09	6.90	88.74
12629	65.11	30.82	27.44	7.55	88.61
18810	1188.71	319.49	906.85	60.20	88.52
18770	1131.01	331.57	887.70	58.94	88.48
19577	494.98	92.34	376.82	38.52	88.48
18891	1378.10	409.91	1823.58	193.46	88.48
7914	1537.05	507.11	1256.25	91.53	88.44

TABLE SF: Timepoint	CAPTOPRIL (s)): 336 hrs		" Ally. Doc		9921-5089W0 Yo. 1798397.1
	MontoxMean -	MonToxSD	ToxiMean	TOXSD .	LDAScore
23859	-10.09	17.25	58.28	12.00	99.70
1522	181.53	92.60	-79.60	14.77	99.40
24668	76.16	46.93	1110.08	605.52	99.10
19287	161.27	40.70	265.00	9.72	98.84
735	125.45	37.81	123.72	0.37	98.50
1348	29.21	17.75	0.23	1.09	98.12
16260	66.53	18.90	82.33	0.49	98.07
24696	69.96	48.08	-11.83	7.33	98.07
826	182.43	53.09	63.46	18.28	97.90
15851	203.67	153.22	36.02	16.84	97.69
19120	32.71	25.65	-8.80	2.19	97.43
1480	253.73	66.75	112.96	18.69	97.43
18659	51.26	29.25	80.08	1.30	97.43
15420	59.26	52.41	-14.25	3.04	97.39
2830	654.10	146.54	933.86	42.78	96.92
2658	891.04	236.04	443.07	41.79	96.88
17937	86.63	42.53	-12.26	22.60	96.83
10108	146.26	35.48	61.17	17.32	96.70
1223	191.43	47.65	289.39	15.78	96.66
16048	28.62	52.28	-8.09	3.07	96.66
10048	26.49	16.73	40.49	1.17	96.62
16944	848.22	191.45	550.56	29.83	96.58
546	173.90	50.96	271.42	13.74	96.53
12819		47.07	239.54		96.53
5735	184.28 56.51	21.12	63.63	3.32 0.57	96.45
				0.69	96.40
7956	27.42	11.15	39.67		
12332	567.22	177.99	328.76	21.43	96.32
18346	273.04	62.53	138.13	21.45	96.32
16425	20.91	30.81	-13.41	3.05	96.23
8426	54.76	23.35	27.91	1.50	96.19
9964	14.13	28.09	41.16	1.54	96.15
15395	797.12	155.79	610.46	12.19	96.02
21458	229.75	104.67	360.26	20.40	95.93
15259	227.63	56.54	154.42	4.47	95.93
11057	33.19	29.92	61.96	3.13	95.89
397	116.22	33.32	88.25	1.44	95.89
20429	108.55	34.07	193.71	39.16	95.85
12333	218.47	76.94	109.09	10.32	95.85
12629	64.58	30.66	134.34	14.35	95.76
20833	1255.87	351.77	1350.26	16.51	95.68
10673	62.50	35.24	48.56	1.93	95.63
9518	29.88	23.15	73.23	8.73	95.59
5630	77.86	48.50	150.84	8.45	95.59
16036	64.27	20.66	37.81	2.46	95.55
23773	214.43	85.23	94.48	13.72	95.51
11817	64.91	20.17	98.04	3.96	95.46
13976	459.29	377.97	122.98	19.90	95.46
14926	95.90	28.06	158.69	14.69	95.46
21633	302.73	119.13	151.23	11.45	95.42
910	58.30	22.93	35.87	1.63	95.42
11203	75.66	25.37	58.05	1.05	95.38
16562	188.78	45.14	125.67	5.83	95.29
6362	63.63	39.69	-15.88	19.50	95.25
23237	89.94	41.57	97.86	1.24	95.25
5384	41.70	44.55	1.71	2.18	95.25
22665	128.49	29.89	159.98	3.47	95.21

		-392	!-		
TABLE 5	F: CAPTOPRIL		Aliy, Do	eket No. 4	4921-5039WO
	11(s): 336 hrs	196			No. 1798397.1
ldendifler		MonToxSD	ToxMean	ToxSD	LDAScore
6581	76.47	26.45	43.08	2.63	95.12
260	417.04	114.29	636.83	72.47	95.12
24814	171.62	33.74	127.03	5.19	95.12
13682	178.68	62.82	61.54	20.17	95.08
15028	346.53	107.46	569.31	126.83	95.08
17439	218.65	47.89	292.03	10.02	94.99
15797	20.60	17.76	-7.25	4.56	94.95
17549	1304.87	367.92	1416.98	24.18	94.95
17923	66.78	23.31	48.46	1.01	94.95
23360	178.74	46.59	218.52	2.99	94.91
20099	81.73	30.45	123.33	3.88	94.91
19327	89.68	29.52	50.76	3.50	94.86
5786	125.64	46.92	48.44	13.78	94.86
9929	531.71	124.53	680.04	14.58	94.82
2831	619.77	172.12	917.69	61.75	94.82
10477	88.24	37.19	51.43	2.96	94.82
21013	217.62	342.33	442.35	95.19	94.82
21651	32.33	26.35	2.01	3.21	94.78
19527	47.24	56.67	112.09	11.81	94.78
1921	178.67	59.48	97.23	10.49	94.73
8988	56.52	38.39	8.71	5.65	94.73
6766	481.24	157.08	653.99	16.43	94.73
18862	47.30	24.54	31.00	1.20	94.69
15470	328.11	71.27	198.29	29.91	94.69
3288	10.56	13.80	41.78	12.82	94.69
23109	2081.66	927.60	2211.62	60.64	94.61
7197	197.55	84.70	290.53	17.88	94.56
7279	201.88	88.54	108.84	9.39	94.52
19581	48.08	32.46	66.08	3.68	94.48

TABLE 50	: CARBOPLAT	IN	' Ally. Doc	ket No. 44	1921-5039WO
	(s): 6 livis				
	MonToxMean		1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 -		
6262	739.41	208.31	1532.53	76.80	99.61
24048	689.96	212.19	1411.28	55.30	99.36
17089	1547.81	626.59	4737.22	696.74	98.93
16081	115.61	379.88	645.20	213.95	98.54
25777	403.77	181.40	901.96	112.12	98.37
4933 7476	134.08 90.92	204.01	525.52 272.77	82.57 23.70	98.20 98.11
15171	241.90	58.14 97.44	403.05	21.96	98.03
24049	1518.47	439.97		123.80	98.03
16080	44.66	251.23	2590.21 378.60	161.27	97.99
10093	342.04	117.93	656.06	50.76	97.86
1069	1820.60	698.77	1513.87	16.71	97.64
25480	92.27	34.63	148.37	3.30	97.43
6647	510.34	159.21	1020.51	105.54	97.39
7247	504.54	113.43	793.61	51.11	97.34
18532	285.39	90.75	560.36	75.34	97.04
4067	123.93	59.96	232.57	36.85	96.92
23449	124.24	104.63	362.40	83.14	96.83
8314	95.89	403.64	202.84	47.87	96.79
14159	87.59		200.11	36.34	96.79
		36.45 131.77	819.56		
18280	584.52			51.87 153.82	96.70
23314	71.21	275.95	495.11		96.70
3816	326.31	77.12	471.75	21.93	96.66
17329	215.35	107.06	443.46	37.86	96.62
5461	193.57	116.65	442.55	77.30	96.57
22501	257.26	67.93	377.18	25.53	96.53
15277	900.30	154.78	1210.05	42.20	96.49
23538	101.09	71.79	256.02	74.72	96.44
8849	222.08	71.82	422.51	61.76	96.36
17779	1949.63	787.42	1501.46	41.41	96.36
23574	2277.50	970.11	1768.74	46.20	96.23
1127	1190.78	434.96	1227.98	24.14	96.14
4154	247.57	87.35	387.82	17.78	95.80
3471	80.41	35.57	126.92	4.34	95.76
22211	768.41	165.76	1038.58	34.89	95.72
21815	248.67	55.97	320.59	9.43	95.67
18597	521.69	163.32	851.48	104.45	95.63
2196	564.46	104.22	713.81	13.18	95.54
13598	350.22	110.18	570.66	47.31	95.46
16895	2022.10	903.47	1796.87	31.07	95.42
6522	571.58	153.62	693.92	15.58	95.42
8652	228.95	114.23	482.75	43.58	95.37
20026	64.36	24.25	120.87	11.23	95.33
15192	169.06	132.74	389.95	89.37	95.29
1622	2067.46	903.52	1573.48	71.56	95.29
3823	526.02	154.25	801.75	71.28	95.20
5989	269.96	71.53	419.90	36.98	95.12
3434	324.23	140.30	584.57	69.17	95.07
3156	1279.17	207.18	1810.46	187.54	95.03
10818	464.74	186.59	181.34	44.19	95.03
5575	86.94	36.56	34.41	8.85	94.99
11174	54.05	51.84	117.20	14.24	94.99
14425	213.74	79.18	349.63	37.59	94.94
16417	72.79	29.59	142.55	15.55	94.90
17771	769.30	306.35	1273.57	156.68	94.86
5208	1055.01	435.10	2101.50	490.32	94.82

TABLE	e: Carboplai		Altity, Dia	ceket No. c	44921-50391\\\
menoi	mt(s); (6 hrs			D)@@_	17/93:353977,5
ldendifie	MonToxMean	DEXCT MOUNT). ToxiMeen	ToxSD	LDAScore
3431	1514.30	636.90	1297.11	34.77	94.82
21462	258.49	59.82	358.44	18.51	94.77
3822	921.02	284.55	1300.83	99.42	94.73
15190	1824.97	1183.78	2141.03	82.58	94.69
19111	1604.90	429.51	2027.13	89.10	94.64
14906	383.54	73.03	535.04	39.09	94.47
13144	-27.27	17.64	-54.47	5.38	94.43
18142	1995.09	839.14	1819.48	43.33	94.39
13634	827.03	282.41	1137.48	107.16	94.34
2350	646.60	103.17	832.07	60.88	94.30
1537	31.35	46.10	59.58	13.80	94.26
13239	110.05	49.12	151.36	7.79	94.17
20864	1582.67	662.32	1722.73	33.76	94.13
12402	540.97	155.59	755.78	32.87	94.13
2424	585.64	132.80	807.06	45.48	94.13
15106	1894.10	713.95	1463.01	37.13	94.09
12569	402.63	145.28	719.92	106.90	94.04
2022	270.40	55.81	378.75	24.92	93.83
13633	310.40	151.71	490.70	80.01	93.79
22197	123.97	52.28	217.39	36.02	93.79
13874	42.03	18.39	51.72	2.31	93.74
3533	212.11	64.89	298.00	11.86	93.74
5985	44.47	68.11	62.07	10.39	93.74
21643	1185.08	370.10	960.79	22.49	93.74
5089	70.77	28.48	123.98	19.48	93.74
17211	1445.11	555.30	1017.49	34.14	93.70
15772	28.16	13.60	23.47	26.74	93.66
25689	1435.28	508.41	1360.96	40.18	93.66
22545	252.03	89.09	444.84	127.61	93.62
11954	3134.66	1692.86	2572.78	105.71	93.57
8634	343.04	96.04	542.29	61.79	93.53
13771	91.36	32.90	152.72	14.45	93.53
14871	688.13	217.01	1066.71	67.36	93.53
25435	69.69	23.87	116.39	11.28	93.49
18076	2539.23	1190.89	2104.90	81.41	93.49
1660	6.16	20.59	24.43	41.71	93.44
20817	1076.80	681.07	1252.44	51.53	93.36
22923	57.85	40.50	3.36	7.47	93.32
20508	16.41	9.30	22.98	0.98	93.27
14304	95.81	30.28	138.83	9.76	93.23
23005	911.26	224.88	1191.71	45.14	93.23
16375	1004.76	291.14	1452.05	94.00	93.23
25754	73.97	20.01	103.72	4.72	93.19
820	2460.91	1164.05	2043.71	81.32	93.19
		1		100=	155.15

TABLE 5H: Timepoint(CEPHALORIDI s): 6, 24 lirs		Alty. Dod	ket No. 44 Doc. N	921-5039WQ o. 1793397.1
ldentifier	NonToxMean	NonToxSD	ToxMean	ToxSD	LDAScore
1698	72.61	72.54	355.15	57.50	98.88
25057	-16.00	13.42	28.61	15.15	98.84
23302	115.35	36.99	250.40	26.19	98.45
25098	44.75	40.56	161.77	39.01	98.45
7022	6.54	19.86	87.07	26.85	98.45
18005	16.30	11.61	71.95	28.31	97.64
16318	111.73	57.76	214.34	48.82	97.59
15849	181.68	73.50	381.77	34.83	97.29
23283	520.93	94.30	765.95	70.15	97.25
651	12.40	11.40	69.75	44.55	97.16
16112	56.89	23.53	146.28	35.70	97.12
25198	33.14	18.60	104.50	28.93	97.12
8879	109.65	35.18	198.92	13.77	97.03
19253	280.76	74.42	450.66	32.07	97.03
15376	140.99	45.51	267.24	27.71	96.90
21038	113.15	45.28	284.43	88.10	96.82
20917	114.76	34.55	219.65	37.89	96.69
650	17.69	13.01	74.86	36.83	96.47
11411	252.17	76.16	422.51	26.01	96.35
343	28.78	32.80	129.29	21.40	96.35
16248	128.59	52.73	293.26	99.85	96.30
20843	165.02	37.03	274.00	60.23	96.30
18995	61.26	23.28	126.99	20.34	96.17
7050	68.24	24.46	116.13	10.21	95.92
20753	128.41	31.77	231.50	37.70	95.83
18084	36.85	19.15	95.80	24.41	95.83
1764	97.37	32.56	181.11	28.78	95.83
22413	73.51	35.54	155.12	24.23	95.74
12162	313.72	69.35	466.55	53.66	95.57
9573	313.16	70.67	449.88	24.42	95.53
5458	521.55	111.52	766.69	82.52	95.49
23889	170.87	60.58	320.41	74.89	95.49
1623	84.54	20.77	130.31	10.23	95.44
19254	240.06	74.71	400.30	44.56	95.31
1628	13.93	10.20	37.44	6.38	95.27
672	-2.45	13.09	33.39	16.62	95.27
1855	13.51	8.02	32.00	4.11	95.18
3900	83.63	36.44	170.88	42.74	95.14
15281	172.53	45.57	282.14	34.03	95.10
1582	16.73	13.23	46.93	7.97	95.10
25589	154.39	37.33	240.89	33.81	95.10
627	63.31	19.85	119.93	25.90	95.06
17434	163.74	49.10	263.86	22.79	95.01
21063	75.00	25.88	144.62	27.06	94.97
14353	61.97	20.12	119.73	30.32	94.93
6850	87.12	29.12	157.80	23.31	94.93
2059	139.66	39.69	243.75	48.13	94.84
16333	132.19	38.88	212.16	19.97	94.71
25377	31.75	20.09	77.51	19.81	94.50
23282	277.48	51.59	385.61	37.11	94.50
12058	109.37	35.93	194.17	43.87	94.45
9952	163.81	35.63	235.14	38.00	94.37
22196	58.71	24.71	123.83	37.19	94.37
16121	110.03	60.28	208.77	16.39	94.33
24640	177.93	59.42	295.24	35.00	94.33
1946	46.78	14.37	67.25	3.13	94.28

Timepoint(s): 6, 24 hrs			Aity. Dodkai No. 44921-5089WC Doc. No. 1793397.1			
ldendifier	NonToxWean	NonToxSD	ToxiMean	ToxSD	LDAScore	
12259	-0.17	9.62	21.16	7.74	94.15	
6790	62.58	24.20	118.33	27.25	94.07	
21802	45.94	21.21	103.13	35.44	94.07	
14125	128.30	50.32	211.96	15.22	94.07	
18183	31.28	14.65	67.29	14.30	94.02	
17225	162.76	46.10	264.51	40.73	93.94	
20514	83.93	25.89	146.04	21.67	93.94	
1342	31.31	15.65	72.08	26.80	93.81	
16616	88.26	37.98	165.12	31.68	93.77	
4386	55.88	26.02	117.38	28.79	93.77	
13464	30.34	17.92	75.47	19.34	93.68	
11358	82.35	33.88	151.50	27.12	93.59	
7866	43.15	16.17	81.51	16.07	93.51	
22967	163.95	54.65	241.59	15.86	93.47	
8385	57.15	28.33	114.53	19.32	93.42	
24748	-23.01	37.80	49.31	11.34	93.38	
16059	50.12	14.86	84.72	17.27	93.38	
16122	117.41	41.42	186.27	14.87	93.34	
1350	143.74	30.94	209.15	28.63	93.34	
8384	39.05	17.03	73.91	10.64	93.29	
18259	216.96	155.49	545.75	138.45	93.28	
20724	48.65	21.46	90.26	16.90	93.25	
352	87.58	53.82	157.16	24.98	93.21	
10740	26.94	24.94	69.61	15.31	93.12	
1394	24.70	11.31	42.23	3.80	93.12	
22466	462.31	92.72	638.40	68.34	92.99	
13684	467.12	135.55	762.39	150.04	92.99	
14768	85.99	47.21	225.50	66.59	92.98	
13285	71.32	19.47	109.95	16.43	92.91	
1537	29.82	40.75	245.05	131.87	92.90	
18442	38.53	17.21	75.30	16.45	92.86	
1183	52.85	29.47	214.38	101.63	92.85	
127	19.05	14.19	48.31	9.93	92.73	
1399	200.20	80.01	449.25	95.96	92.68	
11203	75.19	24.99	127.88	16.58	92.65	
870	22.03	9.87	40.53	4.22	92.61	
4415	38.66	18.55	77.02	14.75	92.61	
373	32.60	56.68	306.93	119.81	92.60	
22524	112.62	47.12	195.09	26.00	92.56	
6951	88.94	34.80	155.83	27.94	92.56	
13023	176.39	400.42	1491.81	754.76	92.55	
17836	102.26	27.02	156.20	28.41	92.48	
7051	62.10	22.11	105.49	19.48	92.48	
18749	115.18	42.04	202.78	33.94	92.39	

TABLE 51:	CIDOFOVIR					
	(s): 120 hrs				lo. 1793 3 97.1	
ldentifier	NonToxMean	NonToxSD	ToxMean	- OSXOT	LDAScore	
18609	203.96	70.15	541.22	19.78	99.70	
20674	83.03	21.84	203.76	22.76	99.61	
4312	76.73	35.26	365.78	59.40	99.57	
24041	12.84	32.48	31.93	0.33	99.53	
5733	10.73	30.54	617.55	122.19	99.44	
2768	1962.14	417.45	933.17	95.15	99.32	
2410	11.13	10.13	64.03	13.02	99.32	
14289	62.53	19.54	95.56	0.76	99.27	
5689	12.82	19.88	85.25	13.41	99.23	
14594	-17.87	24.41	86.07	17.70	99.23	
24000	64.13	33.98	157.59	7.48	99.19	
8027	0.11	26.38	29.41	3.47	99.19	
18322	2666.91	812.51	1165.22	99.03	99.06	
7324	97.12	49.43	271.43	20.08	99.06	
20903	70.06	46.59	288.96	29.37	99.06	
20757	410.75	211.87	1571.32	228.32	98.97	
1599	22.63	26.70	62.22	8.06	98.97	
5183	215.17	70.24	445.78	24.68	98.97	
4856	122.81	48.54	241.75	6.99	98.97	
2655	42.27	39.82	529.96	152.39	98.97	
10167	189.75	100.26	305.25	11.81	98.97	
21275	225.36	80.40	666.87	94.23	98.93	
22722	73.37	39.40	290.14	53.00	98.93	
20082	75.76	32.27	258.55	44.96	98.93	
912	474.15	83.63	764.00	33.56	98.84	
8002	13.99	15.79	47.18	4.09	98.84	
13158	463.87	106.89	360.87	1.60	98.80	
10200	41.90	30.39	92.98	6.46	98.76	
5572	332.45	146.31	611.55	14.40	98.67	
410	1097.88	255.30	605.30	42.31	98.63	
20755	145.36	116.99	986.50	562.94	98.63	
23376	14.61	16.79	37.95	1.52	98.63	
1600	44.38	66.08	114.35	13.26	98.63	
25705	454.53	109.40	1132.56	152.81	98.59	
13609	240.24	60.23	98.11	9.83	98.59	
24219	315.68	82.27	692.16	77.33	98.59	
26184	204.20	70.54	454.10	35.35	98.54	
16081	117.80	381.49	136.30	8.26	98.54	
7660	57.70	87.65	653.31	318.06	98.54	
22681	222.75	194.18	580.89	50.39	98.54	
9215	90.03	33.75	158.07	6.56	98.50	
25699	137.08	68.59	-11.79	8.27	98.50	
4048	-6.15	31.92	56.08	15.06	98.50	
2729	640.63	170.18	328.03	37.57	98.50	
15981	75.77	29.60	148.46	12.15	98.46	
133	-43.58	39.29	55.82	20.14	98.37	
24707	47.32	82.24	10.71	1.68	98.37	
5193	192.02	179.55	250.16	3.55	98.37	
3981	72.52	103.32	368.64	91.18	98.37	
21893	51.03	33.90	172.83	32.83	98.37	
16168	350.38	204.48	829.66	132.95	98.37	
1262	43.72	51.67	193.22	27.67	98.33	
10289	14.87	14.48			98.24	
17161			127.47	59.89		
	1157.45	414.68	2144.64	136.59	98.24	
26150	780.06	534.26	32.24	49.38	98.24	

		*Affy. Doc	ket No. 44	921-50691//0
(s)): 120 hrs			Doc. K	lo. 1793397.1
NonToxMean	NonTox&D	ToxiMean	ToxsD	LDAScore
737.83	188.48	383.68	38.35	98.24
5.98	20.99	59.41	8.18	98.16
113.22	92.96	448.40	70.70	98.16
362.77	114.65	543.47	30.13	98.12
124.48	46.50	204.85	9.09	98.12
6.10	12.64	43.93	7.10	98.07
173.67	81.62	475.77	74.64	98.07
177.77	53.62	345.26	28.00	98.07
555.26	125.28	290.62	37.25	98.07
35.92	17.39	77.13	4.05	98.03
225.07	67.79	427.11	46.05	97.99
49.65	20.05	91.41	3.94	97.99
203.26	65.91	33.57	32.91	97.99
460.95	159.60	865.38	68.10	97.95
22.90	64.75	174.30	49.09	97.90
				97.90
	424.91		134.17	97.90
	176.96	850.57	34.74	97.86
40.69		135.60		97.86
4.99	10.67			97.82
	90.62			97.82
·				97.82
137.15	130.99	496.58	110.00	97.73
107.70				97.73
168.07	57.61		55.59	97.73
3164.11	929.00		124.06	97.73
				97.73
		97.44		97.69
				97.69
				97.69
				97.65
				97.65
				97.65
224.15	69.07	195.44	2.64	97.60
46.63			1.75	97.60
				97.60
		490.19		97.56
				97.56
				97.56
				97.56
				97.56
				97.56
				97.56
38.37	132.57	554.64	221.30	97.52
	(s)): 120 brs NonToxMean 737.83 5.98 113.22 362.77 124.48 6.10 173.67 177.77 555.26 35.92 225.07 49.65 203.26 460.95 22.90 788.37 910.82 519.62 40.69 4.99 382.89 136.50 137.15 107.70 168.07 3164.11 290.06 38.98 420.04 460.82 251.17 308.88 883.32 224.15 46.63 103.14 214.95 238.74 128.75 1289.56 1680.14	NonToxMean NonToxSD 737.83	NonToxMean NonToxSD ToxMean 737.83 188.48 383.68 5.98 20.99 59.41 113.22 92.96 448.40 362.77 114.65 543.47 124.48 46.50 204.85 6.10 12.64 43.93 173.67 81.62 475.77 177.77 53.62 345.26 555.26 125.28 290.62 35.92 17.39 77.13 225.07 67.79 427.11 49.65 20.05 91.41 203.26 65.91 33.57 460.95 159.60 865.38 22.90 64.75 174.30 788.37 204.50 462.53 910.82 424.91 1651.59 519.62 176.96 850.57 40.69 40.63 135.60 4.99 10.67 25.14 382.89 90.62 215.96 137.15 130.99 496.58 107.70 137.96 508.22 168.07 57.61 297.97 3164.11 929.00 1779.72 290.06 65.29 164.79 38.98 26.45 97.44 420.04 95.61 169.12 460.82 148.89 174.40 251.17 124.12 178.93 308.88 74.48 551.57 883.32 205.24 530.31 224.15 69.07 195.44 46.63 26.47 80.60 103.14 35.12 184.15 214.95 93.98 490.19 238.74 78.96 470.27 122.01 67.80 366.63 813.56 118.65 190.91 1680.14 394.54 881.26	NonTexMean NonTexSD TexMean TexSD 737.83

	CISPAN combin 3): 6, 24 hrs	Affy. Docket No. 44921-5089W0 Doc. No. 1798397.			
	NonToxMean	Clesco Track	ToxMean		LDAScore
8990	276.73	75.18	522.76	68.28	96.93
1460	198.36	85.83	320.23	42.90	95.60
16853	67.12	22.69	121.67	21.69	95.08
13239	108.86	47.65	222.86	33.91	94.73
21355	373.47	117.15	627.82	92.43	94.69
6454	238.70	77.20	419.87	47.88	94.69
1247	1313.77	499.48	559.96	117.97	94.56
6506	233.04	59.13	375.55	54.15	94.34
1585	67.19	32.77	140.57	23.52	94.26
1962	33.04	26.70	76.49	11.48	94.17
18433	18.64	45.96	123.81	42.15	94.13
15050	638.79	181.52	459.28	26.53	94.04
17693	1261.20	375.03	651.01	91.79	93.87
4956	78.53	37.78	155.47	25.21	93.65
16233	68.67	88.34	117.42	12.47	93.18
11445	435.55	103.48	642.14	86.21	92.62
8004	125.10	43.16	255.80	43.43	92.37
1811	10.62	25.84	83.04	26.96	92.28
1542	929.09	263.76	560.16	66.12	91.97
16591	151.89	47.44	236.90	29.21	91.84
18694	52.90	48.32	176.24	47.08	91.68
11524	-14.63	24.06	46.43	29.33	91.59
19080	75.03	55.99	212.58	72.04	91.20
20514	83.88	26.10	127.69	12.79	91.19
15701	37.45	16.42	79.72	13.34	91.16
16122	116.73	39.95	217.85	62.09	90.99
2079	303.23	81.83	406.18	33.43	90.98
19327	88.79	28.83	152.22	21.27	90.86
335	95.54	44.88	191.73	23.12	90.86
14003	817.87	211.94	491.17	61.11	90.86
9104	138.43	38.55	221.59	41.76	90.69
25253	291.50	63.92	430.49	54.59	90.69
23322	1169.44	284.23	807.96	73.20	90.67
24696	68.34	46.61	180.46	53.14	90.60
1552	71.02		121.56	15.30	90.54
19120	31.87	83.66 24.91	90.61	26.26	90.54
17411	78.54	53.72	189.16	45.14	90.51
16121	109.00	58.08	258.63	87.07	90.47
1639	96.38	22.51	149.03	17.77	90.47
1622	2080.70	896.54	796.36	153.89	90.43
9882	487.60	181.11	245.48	60.74	90.41
23852	261.73	101.88	487.29	97.17	90.38
13684	465.64	134.08	778.77	108.34	90.38
7857	51.80	45.32	150.23	38.24	90.38
6281	232.74	67.04	343.63	49.82	90.34
15790	45.82	25.45	87.17	18.57	90.24
23884	40.55	32.64	101.67	19.55	90.24
17682	673.97	178.07	455.67	77.33	90.16
16581	39.15	19.04	83.46	15.21	90.10
24390	161.41	112.42	367.16	79.88	90.12
24390 3886	57.66	28.30	124.53		90.08
3666 13682	176.78	61.74	302.10	24.81 50.60	90.08
	1113.30	393.42			
7262			1879.06	245.81	90.00
11954	3158.62	11682.02	952.13	425.90	89.98

TABLE 51: CISPAN combined Timepoint(s): 6, 24 hrs			Aliy. Dockel No. 44921-5039W0 Doc. No. 1793397.1			
ldendifer .	MonToxMenn	MONTOXSID !	ToxiMean	ToxSD	LDAScore	
819	3116.95	1659.72	925.18	255.80	89.90	
16211	2269.84	1132.19	853.70	212.63	89.90	
1521	20.42	50.86	117.89	40.21	89.78	
23125	4625.10	2534.40	1935.15	332.35	89.77	
9109	1022.92	338.20	736.62	48.42	89.77	
10141	53.73	45.97	150.14	52.96	89.69	
152	72.28	26.46	108.76	14.84	89.68	
17154	198.73	57.61	281.13	33.91	89.59	
6362	62.35	38.95	146.74	29.87	89.56	
10540	19.51	18.83	64.08	25.25	89.52	
651	12.58	12.76	33.89	11.18	89.52	
17086	151.38	43.21	214.56	21.12	89.51	
12020	167.06	60.45	260.26	34.23	89.51	
4121	62.05	23.29	113.40	19.65	89.43	
8211	2875.40	1605.55	956.21	247.45	89.38	
20404	53.25	45.78	130.86	32.28	89.35	
7522	31.59	18.74	75.54	18.10	89.30	
20879	87.24	50.13	179.14	40.07	89.26	
17550	1318.67	366.65	860.58	175.55	89.25	
21950	727.03	149.84	534.23	70.97	89.21	
25405	77.77	28.49	139.55	22.89	89.17	
14125	127.72	49.56	229.81	40.28	89.17	
1611	8.22	23.65	52.29	18.59	89.13	
21685	122.94	42.72	207.33	33.76	89.13	
17524	1176.52	284.08	761.61	160.07	89.09	
10611	10.59	27.29	104.61	49.13	89.06	
22849	197.26	50.25	298.83	40.93	89.00	
1608	12.43	29.83	56.08	17.26	89.00	
4312	76.57	37.18	172.28	72.36	88.97	
1396	47.24	18.54	84.98	24.01	88.92	
20871	52.42	25.81	108.86	22.74	88.92	
10344	22.14	21.48	62.29	12.19	88.83	
15587	38.29	19.07	66.30	11.60	88.82	
9096	9634.12	6768.62	2697.48	901.04	88.82	
4290	94.06	28.34	144.64	19.10	88.79	
24143	192.56	96.48	385.57	70.47	88.74	
12174	80.77	29.46	139.07	31.48	88.74	
17336	65.85	21.84	107.99	13.40	88.74	
25257	101.83	34.03	168.25	31.38	88.74	
20350	154.58	49.68	245.00	41.28	88.70	
11335	661.49	162.55	459.38	57.71	88.64	
24146	219.42	50.91	370.23	55.30	88.63	
16254	5.55	11.81	26.92	12.39	88.61	
20876	1683.23	611.09	940.93	123.03	88.60	
9312	31.90	15.15	74.64	21.22	88.58	

14458 36. 22385 106 11731 43. 23745 201 12903 53. 15503 124 8235 43. 16756 177 16119 15. 11967 172 3608 333 18729 19. 5891 -53 2048 31. 4490 76. 1743 29. 1584 162 16137 6.7 23778 68. 23261 156 808 468 1962 33. 21789 34. 23769 -6.2	000 000 000 000 000 000 000 000 000 00	28.40 51.70 24.10 65.06 18.74 37.32 34.77 53.82 15.19 546.07 110.50 29.07 47.69 20.74 72.26	100.33 345.64 393.30 823.60 124.32 319.29 145.95 327.44 84.22 289.45 62.62 174.49 106.02 125.19		9.96 99.87 99.87 99.83 99.83 99.79 99.74 99.74 99.74 99.74 99.70 99.61
22385 106 11731 43. 23745 201 12903 53. 15503 124 8235 43. 16756 177 16119 15. 11967 172 3608 333 18729 19. 5891 -53 2048 31. 4490 76. 1743 29. 1584 162 16137 6.7 23778 68. 23261 156 808 468 1962 33. 21789 34. 23769 -6.2 23070 125 12400 13. 16676 38. 23992 5.3 14430 34. 11969 96. 4312 77. 4967 33. 2079 303 22816	6.64 29 1.94 .64 4.27 .83 7.83 .30 25.18 3.84 .92 3.03 .69 .73 .35 2.19	51.70 24.10 65.06 18.74 37.32 34.77 53.82 15.19 546.07 110.50 29.07 47.69 20.74 72.26	345.64 393.30 823.60 124.32 319.29 145.95 327.44 84.22 289.45 62.62 174.49	8.69 43.69 110.42 6.90 21.43 18.00 3.02 6.17 25.19 5.74 19.16	99.87 99.87 99.83 99.83 99.79 99.79 99.74 99.74 99.74
22385 106 11731 43. 23745 201 12903 53. 15503 124 8235 43. 16756 177 16119 15. 11967 172 3608 333 18729 19. 5891 -53 2048 31. 4490 76. 1743 29. 1584 162 16137 6.7 23778 68. 23261 156 808 468 1962 33. 21789 34. 23769 -6.2 23070 125 12400 13. 16676 38. 23780 25. 25545 94. 11969 96. 4312 77. 4967 33. 2079 303 22816	6.64 29 1.94 .64 4.27 .83 7.83 .30 25.18 3.84 .92 3.03 .69 .73 .35 2.19	51.70 24.10 65.06 18.74 37.32 34.77 53.82 15.19 546.07 110.50 29.07 47.69 20.74 72.26	345.64 393.30 823.60 124.32 319.29 145.95 327.44 84.22 289.45 62.62 174.49	8.69 43.69 110.42 6.90 21.43 18.00 3.02 6.17 25.19 5.74 19.16	99.87 99.87 99.83 99.83 99.79 99.79 99.74 99.74 99.74
11731 43. 23745 201 12903 53. 15503 124 8235 43. 16756 177 16119 15. 11967 172 3608 333 18729 19. 5891 -53 2048 31. 4490 76. 1743 29. 1584 162 16137 6.7 23778 68. 23261 156 808 468 1962 33. 21789 34. 23769 -6.2 23070 125 12400 13. 16676 38. 23992 5.3 14430 34. 11969 96. 4312 77. 4967 33. 2079 303 22816 23. 3609 <	29 1.94 .64 4.27 .83 .7.83 .30 .25.18 .3.84 .92 .0.03 .69 .73 .35 .2.19	24.10 65.06 18.74 37.32 34.77 53.82 15.19 546.07 110.50 29.07 47.69 20.74 72.26	393.30 823.60 124.32 319.29 145.95 327.44 84.22 289.45 62.62 174.49	43.69 110.42 6.90 21.43 18.00 3.02 6.17 25.19 5.74 19.16	99.87 99.83 99.83 99.79 99.79 99.74 99.74 99.74
23745 201 12903 53. 15503 124 8235 43. 16756 177 16119 15. 11967 172 3608 333 18729 19. 5891 -53 2048 31. 4490 76. 1743 29. 1584 162 23778 68. 23261 156 808 468 1962 33. 21789 34. 23769 -6.2 23070 125 12400 13. 16676 38. 23780 25. 25545 94. 11969 96. 4312 77. 4967 33. 2079 303 22816 23. 3609 407 4361 90. 13682 <t< td=""><td>1.94 .64 4.27 .83 7.83 .30 .25.18 3.84 .92 3.03 .69 .73 .35 .2.19</td><td>65.06 18.74 37.32 34.77 53.82 15.19 546.07 110.50 29.07 47.69 20.74 72.26</td><td>823.60 124.32 319.29 145.95 327.44 84.22 289.45 62.62 174.49</td><td>110.42 6.90 21.43 18.00 3.02 6.17 25.19 5.74</td><td>99.83 99.83 99.79 99.79 99.74 99.74 99.74</td></t<>	1.94 .64 4.27 .83 7.83 .30 .25.18 3.84 .92 3.03 .69 .73 .35 .2.19	65.06 18.74 37.32 34.77 53.82 15.19 546.07 110.50 29.07 47.69 20.74 72.26	823.60 124.32 319.29 145.95 327.44 84.22 289.45 62.62 174.49	110.42 6.90 21.43 18.00 3.02 6.17 25.19 5.74	99.83 99.83 99.79 99.79 99.74 99.74 99.74
12903 53. 15503 124 8235 43. 16756 177 16119 15. 11967 172 3608 333 18729 19. 5891 -53 2048 31. 4490 76. 1743 29. 1584 162 16137 6.7 23778 68. 23261 156 808 468 1962 33. 21789 34. 23769 -6.2 23070 125 12400 13. 16676 38. 23780 25. 25545 94. 15254 209 23992 5.3 14430 34. 11969 96. 4312 77. 4967 33. 2079 303 22816	.64 4.27 .83 7.83 .30 25.18 3.84 .92 3.03 .69 .73 .35 2.19	18.74 37.32 34.77 53.82 15.19 546.07 110.50 29.07 47.69 20.74 72.26	124.32 319.29 145.95 327.44 84.22 289.45 62.62 174.49	6.90 21.43 18.00 3.02 6.17 25.19 5.74 19.16	99.83 99.79 99.79 99.74 99.74 99.74
15503 124 8235 43. 16756 177 16119 15. 11967 172 3608 333 18729 19. 5891 -53 2048 31. 4490 76. 1743 29. 1584 162 16137 6.7 23778 68. 23261 156 808 468 1962 33. 23769 -6.2 23070 125 12400 13. 16676 38. 23780 25. 25545 94. 15254 209 23992 5.3 14430 34. 11969 96. 4312 77. 4967 33. 2079 303 22816 23. 3609 407 4361 <t< td=""><td>4.27 .83 .7.83 .30 .25.18 .3.84 .92 .0.03 .69 .73 .35 .2.19</td><td>37.32 34.77 53.82 15.19 546.07 110.50 29.07 47.69 20.74 72.26</td><td>319.29 145.95 327.44 84.22 289.45 62.62 174.49 106.02</td><td>21.43 18.00 3.02 6.17 25.19 5.74 19.16</td><td>99.79 99.79 99.74 99.74 99.74 99.70</td></t<>	4.27 .83 .7.83 .30 .25.18 .3.84 .92 .0.03 .69 .73 .35 .2.19	37.32 34.77 53.82 15.19 546.07 110.50 29.07 47.69 20.74 72.26	319.29 145.95 327.44 84.22 289.45 62.62 174.49 106.02	21.43 18.00 3.02 6.17 25.19 5.74 19.16	99.79 99.79 99.74 99.74 99.74 99.70
8235	.83 7.83 .30 25.18 3.84 .92 3.03 .69 .73 .35 2.19	34.77 53.82 15.19 546.07 110.50 29.07 47.69 20.74 72.26	145.95 327.44 84.22 289.45 62.62 174.49 106.02	18.00 3.02 6.17 25.19 5.74 19.16	99.79 99.74 99.74 99.74 99.70
16756 177 16119 15. 11967 172 3608 333 18729 19. 5891 -53 2048 31. 4490 76. 1743 29. 1584 162 16137 6.7 23778 68. 23261 156 808 468 1962 33. 23769 -6.2 23070 125 12400 13. 16676 38. 23780 25. 25545 94. 15254 209 23992 5.3 14430 34. 11969 96. 4312 77. 4967 33. 2079 303 22816 23. 3609 407 4361 90. 13682 177 15504 <	7.83 30 25.18 3.84 92 3.03 69 73 35 2.19	53.82 15.19 546.07 110.50 29.07 47.69 20.74 72.26	327.44 84.22 289.45 62.62 174.49	3.02 6.17 25.19 5.74 19.16	99.74 99.74 99.74 99.70
16119 15. 11967 172 3608 333 18729 19. 5891 -53 2048 31. 4490 76. 1743 29. 1584 162 16137 6.7 23778 68.9 23261 156 808 468 1962 33. 23769 -6.2 23070 125 12400 13. 16676 38. 23780 25. 25545 94. 15254 209 23992 5.3 14430 34. 11969 96. 4312 77. 4967 33. 2079 303 22816 23. 3609 407 4361 90. 13682 177 15504 129 633 <t< td=""><td>.30 25.18 3.84 .92 5.03 .69 .73 .35 2.19</td><td>15.19 546.07 110.50 29.07 47.69 20.74 72.26</td><td>84.22 289.45 62.62 174.49 106.02</td><td>6.17 25.19 5.74 19.16</td><td>99.74 99.70</td></t<>	.30 25.18 3.84 .92 5.03 .69 .73 .35 2.19	15.19 546.07 110.50 29.07 47.69 20.74 72.26	84.22 289.45 62.62 174.49 106.02	6.17 25.19 5.74 19.16	99.74 99. 7 0
11967 172 3608 333 18729 19. 5891 -53 2048 31. 4490 76. 1743 29. 1584 162 16137 6.7 23778 68.9 23261 156 808 468 1962 33. 23769 -6.2 23070 125 12400 13. 16676 38. 23780 25. 25545 94. 15254 209 23992 5.3 14430 34. 11969 96. 4312 77. 4967 33. 2079 303 22816 23. 3609 407 4361 90. 13682 177 15504 129 633 271 21183 <t< td=""><td>25.18 3.84 .92 3.03 .69 .73 .35 2.19</td><td>546.07 110.50 29.07 47.69 20.74 72.26</td><td>289.45 62.62 174.49 106.02</td><td>25.19 5.74 19.16</td><td>99.74 99.70</td></t<>	25.18 3.84 .92 3.03 .69 .73 .35 2.19	546.07 110.50 29.07 47.69 20.74 72.26	289.45 62.62 174.49 106.02	25.19 5.74 19.16	99.74 99. 7 0
3608 333 18729 19. 5891 -53 2048 31. 4490 76. 1743 29. 1584 162 16137 6.7 23778 68.9 23261 156 808 468 1962 33. 23769 -6.2 23070 125 12400 13. 16676 38. 25545 94. 15254 209 23992 5.3 14430 34. 11969 96. 4312 77. 4967 33. 2079 303 22816 23. 3609 407 4361 90. 15504 129 633 271 21183 22.6 24222 101	3.84 92 3.03 69 73 .35 2.19	110.50 29.07 47.69 20.74 72.26	62.62 174.49 106.02	5.74 19.16	99.70
18729 19. 5891 -53 2048 31. 4490 76. 1743 29. 1584 162 16137 6.7 23778 68.9 23261 156 808 468 1962 33. 23769 -6.2 23070 125 12400 13. 16676 38. 23780 25. 25545 94. 15254 209 23992 5.3 14430 34. 11969 96. 4312 77. 4967 33. 2079 303 22816 23. 3609 407 4361 90. 13682 177 15504 129 633 271 21183 22.6 24222 101	92 3.03 69 .73 .35 2.19	29.07 47.69 20.74 72.26	174.49 106.02	19.16	
2048 31. 4490 76. 1743 29. 1584 162 16137 6.7 23778 68.9 23261 156 808 468 1962 33. 23769 -6.2 23070 125 12400 13. 16676 38. 23780 25. 25545 94. 15254 209 23992 5.3 14430 34. 11969 96. 4312 77. 4967 33. 2079 303 22816 23. 3609 407 4361 90. 13682 177 15504 129 633 271 21183 22.6 24222 101	69 73 35 2.19	20.74 72.26		18.28	
4490 76. 1743 29. 1584 162 16137 6.7 23778 68.9 23261 156 808 468 1962 33. 23769 -6.2 23070 125 12400 13. 16676 38. 23780 25. 25545 94. 15254 209 23992 5.3 14430 34. 11969 96. 4312 77. 4967 33. 2079 303 22816 23. 3609 407 4361 90. 13682 177 15504 129 633 271 21183 22.6 24222 101	.73 .35 2.19	72.26	125.19		99.61
4490 76. 1743 29. 1584 162 16137 6.7 23778 68.9 23261 156 808 468 1962 33. 23769 -6.2 23070 125 12400 13. 16676 38. 23780 25. 25545 94. 15254 209 23992 5.3 14430 34. 11969 96. 4312 77. 4967 33. 2079 303 22816 23. 3609 407 4361 90. 13682 177 15504 129 633 271 21183 22.6 24222 101	.73 .35 2.19			16.43	99.61
1743 29. 1584 162 16137 6.7 23778 68.9 23261 156 808 468 1962 33. 23769 -6.2 23070 125 12400 13. 16676 38. 23780 25. 25545 94. 15254 209 23992 5.3 14430 34. 11969 96. 4312 77. 4967 33. 2079 303 22816 23. 3609 407 4361 90. 13682 177 15504 129 633 271 21183 22.6 24222 101	.35 2.19 75	40.04	462.55	50.86	99.61
1584 162 16137 6.7 23778 68.9 23261 156 808 468 1962 33.2 23769 -6.2 23070 125 12400 13. 16676 38. 23780 25.0 25545 94. 15254 209 23992 5.3 14430 34. 11969 96. 4312 77. 4967 33. 2079 303 22816 23. 3609 407 4361 90. 13682 177 15504 129 633 271 21183 22.6 24222 101	2.19 '5	16.21	72.89	1.22	99.61
23778 68. 23261 156 808 468 1962 33. 21789 34. 23769 -6.2 23070 125 12400 13. 16676 38. 23780 25. 25545 94. 15254 209 23992 5.33 14430 34. 11969 96. 4312 77. 4967 33. 2079 303 22816 23. 3609 407 4361 90. 13682 177 15504 129 633 271 21183 22.6 24222 101		43.70	307.85	8.25	99.57
23261 156 808 468 1962 33. 21789 34. 23769 -6.2 23070 125 12400 13. 16676 38. 23780 25. 25545 94. 15254 209 23992 5.33 14430 34. 11969 96. 4312 77. 4967 33. 2079 303 22816 23. 3609 407 4361 90. 13682 177 15504 129 633 271 21183 22.6 24222 101	.99	397.82	87.49	8.36	99.53
808 468 1962 33. 21789 34. 23769 -6.2 23070 125 12400 13. 16676 38. 23780 25. 25545 94. 15254 209 23992 5.39 14430 34. 11969 96. 4312 77. 4967 33. 2079 303 22816 23. 3609 407 4361 90. 13682 177 15504 129 633 271 21183 22.6 24222 101		34.04	179.58	9.21	99.53
1962 33. 21789 34. 23769 -6.2 23070 125 12400 13. 16676 38. 23780 25. 25545 94. 15254 209 23992 5.39 14430 34. 11969 96. 4312 77. 4967 33. 2079 303 22816 23. 3609 407 4361 90. 13682 177 15504 129 633 271 21183 22.6 24222 101	68.96	389.70	925.80	13.87	99.53
21789 34. 23769 -6.2 23070 125 12400 13. 16676 38. 23780 25. 25545 94. 15254 209 23992 5.39 14430 34. 11969 96. 4312 77. 4967 33. 2079 303 22816 23. 3609 407 4361 90. 13682 177 15504 129 633 271 21183 22.6 24222 101	8.33	143.61	160.94	9.90	99.53
23769 -6.2 23070 125 12400 13. 16676 38. 23780 25. 25545 94. 15254 209 23992 5.39 14430 34. 11969 96.9 4312 77. 4967 33. 2079 303 22816 23. 3609 407 4361 90. 13682 177 15504 129 633 271 21183 22.6 24222 101	.33	26.74	101.07	4.72	99.53
23070 125 12400 13. 16676 38. 23780 25. 25545 94. 15254 209 23992 5.3 14430 34. 11969 96. 4312 77. 4967 33. 2079 303 22816 23. 3609 407 4361 90. 13682 177 15504 129 633 271 21183 22.6 24222 101	67	30.58	177.40	21.84	99.53
12400 13. 16676 38. 23780 25. 25545 94. 15254 209 23992 5.3 14430 34. 11969 96. 4312 77. 4967 33. 2079 303 22816 23. 3609 407 4361 90. 13682 177 15504 129 633 271 21183 22.6 24222 101		8.69	33.99	6.30	99.53
16676 38. 23780 25. 25545 94. 15254 209 23992 5.3 14430 34. 11969 96. 4312 77. 4967 33. 2079 303 22816 23. 3609 407 4361 90. 13682 177 15504 129 633 271 21183 22.6 24222 101	5.37	31.23	218.44	4.95	99.49
23780 25.4 25545 94.1 15254 209 23992 5.3 14430 34.1 11969 96.6 4312 77. 4967 33.2 2079 303 22816 23.3 3609 407 4361 90.1 13682 177 15504 129 633 271 21183 22.6 24222 101		10.48	74.60	11.18	99.49
25545 94.3 15254 209 23992 5.33 14430 34.3 11969 96.8 4312 77.4 4967 33.3 2079 303 22816 23.3 3609 407 4361 90.1 13682 177 15504 129 633 271 21183 22.6 24222 101		26.96	110.79	5.32	99.49
15254 209 23992 5.39 14430 34. 11969 96.9 4312 77. 4967 33. 2079 303 22816 23. 3609 407 4361 90. 13682 177 15504 129 633 271 21183 22.6 24222 101		35.93	95.20	12.81	99.44
23992 5.39 14430 34. 11969 96.8 4312 77. 4967 33.4 2079 30.3 22816 23. 3609 40.7 4361 90. 13682 17.7 15504 129 633 27.1 21183 22.6 24222 10.1		48.16	306.94	34.66	99.44
14430 34. 11969 96. 4312 77. 4967 33. 2079 303 22816 23. 3609 407 4361 90. 13682 177 15504 129 633 271 21183 22.6 24222 101		67.16	447.21	21.22	99.44
11969 96.8 4312 77.7 4967 33.4 2079 30.3 22816 23.3 3609 40.7 4361 90.1 13682 17.7 15504 12.9 633 27.1 21183 22.6 24222 10.1		7.25	34.45	3.17	99.44
4312 77. 4967 33.4 2079 303 22816 23.3 3609 407 4361 90. 13682 177 15504 129 633 271 21183 22.6 24222 101		31.78	125.29	8.54	99.40
4967 33.4 2079 303 22816 23.3 3609 407 4361 90.3 13682 177 15504 129 633 271 21183 22.6 24222 101		38.48	303.71	38.83	99.40
2079 303 22816 23. 3609 407 4361 90. 13682 177 15504 129 633 271 21183 22.6 24222 101		37.99	241.97	24.41	99.40
22816 23. 3609 407 4361 90. 13682 177 15504 129 633 271 21183 22.6 24222 101		19.94	94.30	3.72	99.36
3609 407 4361 90. 13682 177 15504 129 633 271 21183 22.6 24222 101		81.26	527.92	20.78	99.32
4361 90. 13682 177 15504 129 633 271 21183 22.6 24222 101		15.17	89.33	8.69	99.32
13682 177 15504 129 633 271 21183 22.6 24222 101		148.70	67.97	9.33	99.32
15504 129 633 271 21183 22.6 24222 101		30.77	193.28	10.19	99.32
633 271 21183 22.6 24222 101		62.54	333.83	4.20	99.32
21183 22.6 24222 101		50.62	447.33	63.50	99.27
24222 101		101.41	110.07	15.08	99.27
		26.03	134.40	19.53	99.27
გე4გ I63.0		52.50	460.48	61.63	99.27
7/20/		30.22	6.09	1.52	99.27
	9.26	35.74	248.67	13.48	99.27
18442 38.6		17.15	99.87	5.14	99.27
14370 32.5 21500 80.3		45.58 76.60	189.80 471.24	11.25	99.23
		76.69		73.44	99.23
	3.45	87.73	552.73	49.74	99.23
	J (17)	219.52	248.56	33.80	99.23 99.23
	3.02	36.00	242.62	24.88	
	31	23.52	203.14	12.02	99.23
	31 0.74	25.98	111.75	3.84	99.23
20890 101 712 1.36	31 0.74 72	46.35 9.08	342.52 60.25	50.55 14.12	99.19 99.19

	CISPLATIN		- Ally, Dod	ket No. 44	921-503
	(s): 163 hrs				<u>lo. 1798</u> 9
ldentifier	NonToxMean	NonToxSD	ToxiMean	ToxSD	LD/AS@
18553	54.23	33.42	166.80	12.30	99.19
15884	183.00	57.89	383.85	23.01	99.19
19722	168.71	55.03	336.25	15.95	99.19
17481	26.11	28.66	137.39	13.89	99.19
5733	12.25	46.24	172.15	34.99	99.19
4895	232.18	95.74	44.77	8.48	99.14
15151	153.83	39.15	302.83	29.57	99.14
14759	21.72	15.86	78.25	9.50	99.14
15039	272.18	79.33	101.53	6.80	99.14
12782	20.00	43.24	135.93	6.16	99.14
23121	19.80	13.92	60.40	2.66	99.14
26292	22.82	12.83	85.01	11.49	99.14
2154	59.03	121.05	244.16	33.75	99.14
21583	125.87	40.62	270.04	27.65	99.14
3006	26.57	35.44	81.75	9.18	99.10
1203	5.08	18.58	86.62	14.26	99.10
24472	234.56	47.65	380.85	13.16	99.10
5729	87.49	38.35	274.95	40.90	99.10
132	-24.93	23.56	50.93	12.76	99.10
1801	97.64	29.41	197.50	14.44	99.10
1993	30.09	21.86	117.89	9.03	99.10
16675	33.74	34.04	112.81	11.29	99.10
1382	57.80	21.22	127.59	5.60	99.10
17586	115.18	35.27	246.61	21.48	99.06
21666	22.81	17.92	87.38	6.42	99.06
2125	76.55	80.69	345.60	72.31	99.06
21709	142.70	29.95	215.46	6.19	99.06
16538	132.96	36.43	245.73	9.25	99.06
2845	667.61	123.51	1098.14	42.15	99.02
753	42.04	16.33	120.03	23.17	99.02
21893	50.92	33.48	203.93	25.50	99.02
21836	29.24	16.81	91.95	8.44	99.02
21817	10.63	13.30	59.28	7.00	99.02
6517	231.85	156.64	721.49	128.57	99.02
1588	61.92	22.15	122.71	6.97	99.02
14564	48.33	23.27	104.89	3.20	99.02
3079	34.64	46.77	152.78	26.18	98.97
7602	193.69	43.89	364.34	31.12	98.97
20816	393.33	173.56	774.42	37.69	98.97
6322	18.88	17.54	122.95	22.89	98.97
17337	510.70	139.07	253.28	6.67	98.97
18161	114.11	43.86	244.42	22.79	98.97
4057	64.54	20.73	139.42	12.02	98.97
22552	314.43	92.85	696.18	70.05	98.93

			- Ally. Doc	ket No. 44921-5039WC		
	(s): 6, 24 hrs NonToxWean	Maniaman	TowWoon!		10. 1798397. 11 DASeers	
20082	75.47	31.84	228.60	43.28	98.84	
1598	47.75	56.92	133.89	25.63	98.58	
15313	11.08	24.34	53.59	5.89	98.54 98.20	
2655	43.13	48.31	177.30	66.12	97.85	
14424	66.72	130.96	272.03 63.70	63.86 27.83	97.85	
17314 21275	5.89 225.06	11.89 80.71	528.11	144.80	97.68	
4047	85.85	50.33	190.04	23.02	97.60	
20116	9.55	31.45	124.39	50.70	97.38	
15382	122.93	203.50	341.43	85.17	97.34	
1521	20.82	51.01	148.25	24.41	97.30	
24146	220.17	51.83	396.31	40.61	97.30	
8990	278.10	77.14	540.97	69.47	97.21	
1884	178.33	37.79	272.14	18.27	97.17	
4933	134.55	205.15	335.87	89.54	97.04	
6506	233.65	59.50	415.92	45.52	97.04	
4944	112.31	56.05	264.36	39.37	96.95	
8004	125.79	44.06	271.11	40.68	96.95	
1993	30.02	21.93	91.88	19.85	96.91	
20506	21.57	7.97	46.99	7.30	96.82	
21462	257.92	58.72	424.63	52.75	96.78	
6974	129.50	46.90	236.12	28.32	96.78	
11549	243.07	64.12	425.53	38.72	96.74	
2905	245.79	107.78	488.69	56.33	96.74	
1811	10.99	26.26	94.19	25.01	96.70	
10839	313.60	67.83	533.54	83.77	96.70	
2468	252.35	62.46	433.73	70.78	96.61	
373	33.94	61.41	162.88	51.32	96.57	
19040	186.25	107.25	258.19	19.20	96.57	
15299	87.98	61.10	206.62	59.64	96.57	
13684	467.21	135.45	831.58	69.76	96.52	
910	57.76	21.93	136.53	45.93	96.35	
4477	11.91	8.60	37.00	7.79	96.27	
20871	52.68	25.99	122.97	14.46	96.22	
16853	67.36	22.84	136.28	21.27	96.18	
23473	156.94	56.94	338.21	73.37	96.18	
2536	393.91	124.60	585.25	20.89	96.14	
10015	232.29	77.80	340.64	28.00	96.14	
18694	53.51	48.91	198.02	43.87	96.09	
16284	40.93	27.58	129.79	27.84	96.09	
18375	121.97	28.16	198.57	23.40	96.05	
23314	70.46	275.32	498.55	211.58	96.05	
651	12.67	12.78	40.79	8.43	95.97	
3266	133.11	36.44	247.88	45.93	95.97	
1460	198.97	85.95	340.15	49.64	95.97	
20065	86.81	30.90	170.50	27.85	95.97	
15301	38.37	68.10	96.58	21.67	95.92	
23448	169.78	101.84	327.19	44.52	95.92	
15003	36.27	96.97	63.65	10.34	95.88	
26184	203.75	69.61	423.29	118.85	95.88	
8336	26.96	37.28	139.71	61.60	95.79	
6362	62.77	39.27	160.35	23.77	95.75	
14003	815.94	212.81	484.61	51.19	95.67	
9339	357.63	88.22	598.27	74.10	95.62	
6384	60.88	59.65	127.67	27.22	95.54	
15345	201.66	67.69	322.91	40.01	95.45	

100	: CISPLATIN		Ally. Do		4921-5009W
	((S)): 6, 24 hrs				No. 1798397.
ldentifier	NonToxMean	NonToxSD	ToxiMean	Toksd,	LDASCOR
22849	197.75	50.59	318.11	30.11	95.41
23868	178.04	289.84	277.77	47.26	95.41
16233	68.93	88.14	123.52	11.88	95.36
6454	239.80	78.37	418.57	35.68	95.36
21061	58.86	27.77	128.21	17.87	95.36
24143	193.56	97.20	413.12	53.39	95.32
15296	137.87	56.84	269.21	33.23	95.32
22374	148.45	41.20	247.50	28.04	95.28
13239	109.46	48.17	236.67	36.75	95.28
8768	64.97	26.90	141.33	30.56	95.28
1542	927.15	264.15	512.83	50.15	95.28
22352	156.66	103.69	254.35	25.00	95.24
14051	133.17	35.29	219.32	28.80	95.24
9343	189.65	67.21	349.25	44.44	95.24
1247	1309.30	501.39	550.55	103.22	95.19
7857	52.35	45.86	156.83	28.07	95.19
11727	230.50	78.02	409.08	59.02	95.19
1639	96.67	22.79	153.44	17.91	95.11
15374	138.74	36.97	231.57	31.99	95.02
3899	141.45	61.33	242.19	14.71	94.98
25405	78.08	28.73	149.92	18.38	94.98
23872	49.59	92.93	101.10	24.53	94.98
24368	244.08	80.76	439.99	52.18	94.98
10818	465.48	185.96	140.29	49.38	94.98
17693	1257.86	376.37	596.02	96.76	94.98
25253	292.22	64.49	449.51	58.18	94.89
11708	319.92	92.39	486.60	46.21	94.89
17908	63.14	60.90	124.23	27.40	94.89
24028	407.75	96.21	644.04	67.17	94.85
11455	115.78	48.10	202.56	32.14	94.85
20870	19.10	29.21	92.31	26.59	94.81
3931	94.55	29.06	155.45	15.94	94.68
6581	76.02	26.19	132.63	10.11	94.68
1447	208.86	37.67	272.57	11.93	94.64
22501	257.01	67.60	383.79	63.10	94.64
10720	153.08	43.55	217.97	97.98	94.64
20591	25.22	22.60	68.79	11.62	94.59
1292	62.12	24.60	116.95	15.58	94.55
21355	374.95	118.50	635.77	102.99	94.51
23852	262.89	102.86	519.85	100.21	94.51
18689	360.99	81.71	541.28	55.93	94.51
4426	222.88	38.63	305.45	29.97	94.51
23563	24.57	33.24	96.80	28.33	94.46

TABLE 5M:	: CISPLATIN 3): 6, 24, 163 hrs		Ally. Docke	No. 44921 Dec. 1	1-5039VVO Vo. 1798397.
i nors Sannali	· · · · · · · · · · · · · · · · · · ·		ToxWean *		LDAScore
1521	20.35	50.42	152.11	27.16	97.37
1884	178.00	37.40	272.66	21.76	97.03
16284	40.57	26.94	134.66	25.22	96.99
18694	52.96	48.08	203.55	35.96	96.77
	312.93	66.86	524.22	78.99	96.77
10839 11549	242.48	63.43	420.82	36.27	96.77
8990				72.77	
6362	277.44 62.37	76.38	515.35 167.05	23.05	96.60 96.60
	10.74	38.73	89.58	21.97	96.60
1811		25.96			
4477	11.83	8.50	36.45	6.35	96.55
4047	85.57	50.18	181.03	26.48	96.55
16853	67.12	22.50	136.44	18.16	96.55
3266	132.70	35.81	249.33	37.38	96.47
13684	466.14	134.44	811.93	63.86	96.47
2905	244.60	105.83	525.71	99.02	96.43
1460	198.53	85.75	335.50	43.05	96.38
23314	69.18	274.82	477.84	185.93	96.25
17693	1260.38	374.54	570.60	88.76	96.21
15301	38.18	68.12	96.20	20.88	96.12
17894	46.77	18.63	103.03	28.25	96.08
10015	231.46	76.52	389.17	82.89	95.99
19040	185.68	106.97	292.78	52.89	95.95
18375	121.75	27.96	193.24	20.67	95.74
4426	222.53	38.25	311.48	25.65	95.74
7857	51.97	45.46	159.10	25.13	95.65
6454	239.16	77.72	421.14	34.72	95.65
15296	137.17	55.52	295.99	61.04	95.56
10818	467.00	184.46	97.87	70.28	95.52
23852	261.84	101.45	536.68	85.55	95.52
1542	928.65	263.36	505.79	51.94	95.48
20090	122.37	27.65	178.94	11.38	95.31
1247	1312.52	499.20	486.71	136.84	95.31
13682	176.87	61.56	326.18	46.83	95.26
14003	817.40	211.66	451.12	81.85	95.26
12478	82.11	29.97	149.88	36.04	95.22
1585	67.31	32.78	148.67	23.18	95.22
9339	357.01	87.72	575.84	68.74	95.13
23868	177.77	290.29	269.73	44.26	95.09
20591	25.06	22.48	69.25	11.11	95.05
3352			75 7 .04		
	439.29	115.94		153.95	95.00
6974	129.29	46.84	219.21	32.67	94.88
8888	58.70	29.09	133.86	26.30	94.88
21061	58.67	27.62	123.00	16.55	94.83
7262	1114.68	393.99	1942.97	210.00	94.83
24368	243.52	80.33	427.84	45.50	94.70
25253	291.81	64.18	436.10	53.03	94.53
20921	26.87	19.97	73.82	15.18	94.44
15438	65.27	31.97	154.80	42.05	94.36
12174	80.78	29.36	153.45	25.53	94.32
808	469.71	142.88	210.55	55.06	94.23
19667	34.63	16.92	74.26	11.91	94.19
3886	57.79	28.39	128.86	25.08	94.19
20082	75.08	31.10	214.13	47.37	94.08
14051	132.96	35.15	209.45	29.46	94.01
5073	188.58	69.99	346.14	69.21	93.93

TABLES	in: Cisplatin	W 4000	Alty. Docke	1 No. 4492	1-5039WO
Timepoint(s): 6, 24, 163 hrs					No. 1793397.1
ldentifier					LDAScore
10344	22.23	21.56	63.33	9.18	93.84
15386	93.23	133.87	380.59	101.27	93.76
9882	486.99	181.36	244.23	35.18	93.76
4443	259.36	71.87	429.91	75.00	93.76
16080	44.85	252.23	176.73	162.58	93.76
22005	63.42	50.09	187.23	39.58	93.71
15313	10.94	24.26	52.72	7.91	93.65
6384	60.76	59.71	117.04	26.05	93.63
15701	37.60	16.66	75.18	12.81	93.58
22257	33.56	15.61	61.03	7.13	93.54
2655	42.88	48.17	155.13	62.85	93.52
4198	698.17	161.70	448.59	56.51	93.50
6522	569.21	150.95	877.20	111.47	93.28
19128	112.63	38.66	180.82	19.01	93.15
17314	5.74	11.53	57.89	29.34	93.13
22871	101.59	29.83	165.82	22.84	93.11
21275	224.35	79.76	492.77	134.59	93.09
1727	43.38	60.84	96.90	23.97	93.07
19249	264.62	60.02	356.76	20.60	93.07
1993	29.72	21.35	101.33	20.81	93.04
4584	82.33	29.05	125.60	12.25	92.98
24162	523.84	133.30	788.13	85.20	92.94
7522	31.71	18.92	74.24	15.00	92.89
17713	134.82	29.04	191.13	21.86	92.85
1428	-7.27	15.96	40.03	16.58	92.83
14776	99.41	35.18	153.67	26.50	92.81
3418	333.44	86.68	495.74	46.58	92.81
4199	529.94	132.34	301.96	61.95	92.64
21685	123.15	42.92	207.52	32.56	92.64
7023	363.44	76.61	483.68	29.43	92.55
4420	40.76	32.16	92.81	27.98	92.48
4121	62.21	23.49	110.83	18.20	92.42
18995	61.31	23.53	109.28	20.56	92.42
14665	151.38	37.30	219.57	23.41	92.33
11404	134.16	54.82	304.44	53.63	92.14
910	57.47	21.40	138.17	36.81	92.10
24081	117.08	63.79	235.77	50.20	92.03
22351	45.44	32.53	86.73	20.28	91.99
16012	72.47	31.00	153.59	33.17	91.97
22211	766.80	164.47	1061.59	90.34	91.95
727	210.80	38.94	274.04	15.91	91.90
9104	138.35	38.03	253.38	40.14	91.88
10417	43.82	25.97	125.71	32.61	91.88
10611	10.74	27.42	115.20	49.23	91.84
1314	262.03	49.07	394.88	48.07	91.79

		-407	'-				
	: CITRININ : (s): 6, 24 lirs		Aiiy. De		1921-50:9W(No. 1793397.		
	NonToxMean	Manifor (SD)	Mass Margin				
17541	622.47	209.71	2524.98	697.44	98.92		
6108	533.46	112.70	1024.58	102.59	98.62		
25064	962.08	317.37	2651.31	379.95	98.58		
1698	70.27	55.64	598.98	232.89	98.49		
8820	130.72	105.90	755.03	221.60	98.41		
23917	725.54	174.69	1782.62	536.25	98.36		
20817	1043.62	545.12	5020.35	2088.07	98.32		
15391	756.64	170.28	1510.23	272.65	98.24		
20864	1562.02	620.45	4051.37	596.98	98.19		
24192	70.11	37.23	212.99	53.54	97.93		
20818	665.29	354.08	2965.76	1254.63	97.93		
1340	192.34	49.85	114.30	11.66	97.76		
20035	180.50	107.18	446.34	61.98	97.72		
25525	1057.73	339.31	2228.58	326.32	97.55		
18989	782.09	261.73	1560.14	205.30	97.46		
3431	1496.13	608.90	3517.88	499.00	97.42		
13723	734.46	282.33	1643.16	339.11	97.29		
353	173.42	82.57	323.18	43.00	97.25		
15848	1318.65	418.94	2622.73	442.07	97.16		
634	1135.42	374.65	2281.52	441.32	96.86		
354	214.25	93.76	406.34	75.12	96.77		
7681	101.15	43.91	212.89	41.46	96.64		
13610	357.22	70.16	213.99	29.24	96.56		
5601	970.57	259.22	473.69	102.75	96.47		
3876	30.44	14.34	1.17	8.63	96.13		
24375	115.03	40.98	208.88	57.88	96.04		
8212	2232.57	1104.98	5289.94	717.26	95.96		
15106	1878.08 1214.56	698.66	3540.36	359.29	95.74		
14670 15189	1735.59	325.29 1153.61	1917.56 4557.38	318.04 1226.17	95.70 95.70		
24496	122.73	40.31	47.99	15.89	95.57		
20895	332.13	100.81	137.61	37.56	95.52		
20876	1661.97	599.79	3113.86	382.39	95.52		
18533	35.68	17.84	5.09	4.52	95.52		
6630	1393.25	256.89	913.59	112.68	95.48		
20844	837.88	286.57	1606.81	216.13	95.44		
15850	1209.74	343.26	2046.05	316.40	95.44		
7315	-13.72	19.65	37.49	20.35	95.44		
9254	247.89	49.11	157.69	17.49	95.44		
15363	443.12	132.07	716.96	123.62	95.31		
18359	245.00	113.72	546.56	83.87	95.22		
8211	2822.70	1582.68	6312.46	772.17	95.22		
16831	42.34	13.79	13.06	7.30	95.22		
18644	2209.11	1246.61	5017.03	698.10	95.14		
15190	1803.71	1156.33	4453.59	1177.82	95.09		
15201	1480.71	527.22	2864.67	616.92	95.09		
18205	277.90	58.81	408.93	34.56	95.09		
19094	1054.85	243.62	1623.72	219.27	95.05		
17108	219.63	44.77	132.90	13.45	95.01		
10464	131.59	35.74	68.61	13.81	94.84		
9409	102.66	32.91	171.76	24.78	94.84		
7586	795.05	201.44	401.14	109.74	94.79		
18800	2701.84	1144.69	5340.05	791.36	94.79		
25675	688.20	206.46	1130.86	172.21	94.75		
2697	1222.06	354.81	1908.51	176.60	94.75		
10267	2101.59	872.11	4127.29	650.25	94.71		

TAN E SIN		initial was in the	AYYY. Die	ශ්යා No. W	1921-50000000
	(s):: 6, 24 hrs		rasji Ze		lo. 1798397.1
ldendifier	NonToxMean	NonTox&D.			
22773	230.97	52.31	131.59	26.18	94.54
1651	880.31	240.55	631.98	31.80	94.19
17494	219.24	41.89	138.20	23.13	94.15
244	51.19	35.18	14.03	8.20	94.15
17693	1247.25	372.94	2027.18	231.36	94.10
6946	389.05	103.41	200.78	37.64	94.02
23783	436.13	76.56	298.50	28.57	93.93
19408	1997.30	674.69	2937.88	154.93	93.89
20088	383.97	79.45	244.56	32.48	93.89
16272	192.25	63.76	102.37	22.94	93.89
2866	642.47	211.99	276.00	84.07	93.85
16954	48.70	79.17	202.92	44.32	93.80
21685	124.50	43.36	59.69	10.37	93.72
1719	145.21	38.11	80.93	12.30	93.67
20810	1256.69	398.25	2088.75	313.45	93.63
5049	298.40	65.65	175.28	28.59	93.63
1814	172.31	47.40	99.15	13.17	93.63
16193	101.42	30.67	44.23	15.12	93.59
15017	1007.41	395.69	2150.20	484.37	93.58
17686	1014.59	265.55	1558.32	151.72	93.50
20803	432.89	100.25	912.31	140.76	93.49
1537	29.21	35.65	294.59	185.42	93.45
1399	198.89	71.28	576.63	288.99	93.45
22583	26.67	14.48	3.43	7.60	93.37
3091	784.77	186.23	457.77	105.80	93.37
9029	430.46	93.52	614.72	63.36	93.37
16849	114.53	44.11	44.12	13.14	93.33
22414	58.52	33.14	101.78	22.01	93.33
8283	122.89	43.92	338.56	122.17	93.32
20918	440.21	126.29	269.42	29.15	93.29
25069	131.62	55.06	390.27	136.77	93.28
19067	175.20	51.04	88.87	21.48	93.24
7022	6.12	16.94	127.93	68.30	93.24
723	32.80	15.28	8.05	5.98	93.20
2242	2295.34	607.45	1325.08	393.96	93.16
24390	165.35	113.57	-9.44	39.74	93.07
17211	1434.51	548.50	2462.85	274.87	93.03
22406	79.97	30.24	31.26	10.08	92.90
24469	1169.56	333.98	1827.58	165.96	92.86

The second secon	: COLCHICINE (S): 6, 24, 43 hrs	3 35.			921-5039WC o. 1793397.1
ldendifer	NonToxMean	NonToxSD	ToxiMean ::		LDAScore
23166	132.81	58.53	371.64	112.54	97.29
4412	377.59	61.12	558.75	77.91	96.99
18151	1133.81	261.37	563.27	117.10	96.47
15964	1187.13	325.69	508.38	159.50	96.12
11618	419.37	129.59	119.80	80.64	96.12
16882	177.88	54.70	82.96	14.87	95.78
24321	722.16	202.93	318.04	123.60	95.43
9097	258.85	82.10	115.52	31.47	95.05
20001	1683.96	373.32	1059.37	114.06	94.92
16913	1386.59	297.09	894.57	81.02	94.75
17887	1461.63	317.71	838.24	208.19	94.70
16924	445.74	155.90	188.36	46.16	94.57
20988	1221.44	211.78	853.97	84.52	94.49
22271	275.62	66.08	156.87	37.40	94.32
2222	802.84	200.86	537.76	35.92	94.06
6806	1236.90	342.77	566.10	189.93	94.01
13855	22.06	28.83	139.05	42.55	93.95
20513	57.97	25.96	166.84	71.44	93.82
9296	1306.96	257.77	840.48	105.26	93.80
643	62.16	32.97	14.32	5.63	93.71
16982	128.17	257.40	1462.72	552.54	93.60
20312	405.72	95.10	209.02	92.75	93.58
4073	562.17	184.40	230.54	83.32	93.58
3925	477.10	1119.06	265.65	66.28	93.45
2913	736.68	160.48	481.77	40.00	93.24
10984	2020.41		1032.75	I	92.98
22321	100.73	564.71 64.58	321.87	348.86	
2767	44.79		182.37	131.71	92.96
4151	563.27	40.13 167.25		66.23 111.75	92.91 92.89
7615	90.46	40.50	920.16		
			169.26	27.71	92.81
17399 6552	1980.57	449.65	1255.63	187.29	92.64
13111	1594.49 228.20	296.14 73.35	1077.60 132.80	126.50	92.64
				17.16	92.59
13727 18642	124.04	53.37	38.71	19.99	92.55
3050	974.76	203.55	664.60	62.35	92.55
3050 3438	91.21	49.08	313.78	134.78	92.53
	114.03	57.15	38.70	14.25	92.51
20405	33.56	28.74	132.47	64.39	92.48
14185	202.17	89.20	491.51	237.69	92.48
16849	114.55	44.14	48.23	13.11	92.46
12901	1626.70	415.03	1002.61	177.27	92.42
20697	1421.26	277.19	931.95	163.03	92.25
3837	359.37	91.36	235.96	24.61	92.25
17361	145.97	59.92	56.10	14.57	92.20
17329	213.41	103.82	524.01	86.38	92.18
15600	774.27	210.27	435.87	107.62	92.08
16879	1207.58	276.77	796.63	98.39	92.03
1330	500.26	160.51	241.64	84.25	92.03
22152	7.72	33.71	81.27	38.69	91.79
21053	105.50	59.69	39.98	16.30	91.69
7324	360.96	79.01	153.55	65.98	91.58
7540	155.81	94.07	328.58	92.44	91.58
16128	286.68	62.49	192.02	31.18	91.56
4790	165.31	84.94	49.50	20.48	91.52
23115	570.61	171.16	295.42	91.28	91.47

	0: COLCHICINE 14(s): 6, 24, 43 his		Ally, Doo		1921-5039VVQ 10. 1793397.1
			ls		
	NonToxMean.	de la company de		HALL SOMETHING THE PROPERTY OF THE PARTY OF	
11057	32.46	28.53	120.72	43.20	91.41
7537	230.23	69.42	125.41	29.85	91.34
19822	1596.44	410.54	942.00	210.50	91.26
17386	303.88	93.74	128.16	41.06	91.02
17248	2568.34	534.75	1704.94	295.46	91.00
15191	2013.99	1219.17	2599.48	219.23	90.96
1141	240.56	63.32	390.27	85.33	90.93
3099	966.22	189.19	660.01	95.36	90.70
21024	577.99	113.27	331.56	66.79	90.59
8709	148.12	48.57	78.15	19.09	90.57
19731	226.36	215.21	98.99	29.40	90.57
6250	492.94	104.90	372.14	23.59	90.53
117	21.16	17.56	-5.10	9.54	90.44
17401	907.31	422.79	1550.66	328.51	90.42
15377	25.87	15.51	55.07	8.41	90.42
17326	22.55	24.06	93.51	28.61	90.42
22697	59.78	40.38	12.83	13.23	90.35
14595	87.77	36.04	175.49	38.60	90.29
9223	150.65	64.81	54.19	23.82	90.27
8785	209.90	55.55	288.76	24.92	90.22
9339	360.32	89.43	226.70	47.31	
23253	624.10	163.49	380.42	78.17	90.09
25907	19.74	25.13	47.86	14.93	90.09
15893	1733.99	343.83	1225.85	165.26	90.05
23514	407.55	167.53	162.38	70.14	89.97
3875	510.89	136.65	232.91	81.51	89.90
406	362.17	77.76	250.18	48.87	89.88
18343	437.98	107.72	280.49	46.68	89.79
25461	49.71	22.92	20.96	5.68	89.75
10789	326.18	107.75	117.23	67.68	89.73
23145	44.09	20.22	87.09	15.12	89.73
4048	-7.14 53.20	26.09	121.21 171.90	153.77	89.66
11174	- "	50.67		46.40	89.64 89.45
23709	2491.22 194.78	1205.88	2706.26	145.13	1
23224		48.70	129.40	19.04	89.41
11215	143.29	71.54	28.34 134.33	56.76	89.23
19479	276.88	106.02		36.73	89.23
15872	152.44	63.81	467.24	163.94	89.23
10985	1146.00	265.29	633.75	164.98	89.17
18451	1444.75	418.00	943.24	132.82	89.10
812	157.19	35.17	88.26	32.94	89.04

PCT/US02/16173

	CYCLOPHOSP!		Ally, l		4921-5009WO
	XON OXXIIDEN S)): O Wis	<u>פאפיניסוויסאון</u>	TOXIME an	Thoms how	No. 1798897.1
lcenumer 17089	11545.78	604.39	6128.97	335.19	199.53
16081	113.02	372.25	1532.22	184.12	99.40
23619	327.35	96.06	660.24	21.96	99.32
5393	-31.45	22.77	27.35	5.29	99.32
24049	1517.02	432.93	3280.45	348.50	99.10
22698	261.92	112.70	-415.61	188.19	99.06
26222	371.68	149.80	1265.79	372.61	98.76
18118	895.16	271.71	2268.31	465.94	98.67
16469	1143.78	291.23	1083.34	3.98	98.67
17066	37.10	19.62	-6.41	5.49	98.67
7084	123.08	117.51	622.25	118.49	98.59
24213	1604.12	394.78	3341.49	503.25	98.54
3470	143.86	59.86	458.18	196.11	98.54
23711	4518.75	2192.54	16927.73	3240.54	98.50
18831	4165.13	1320.79	10676.69	1466.84	98.50
10831	289.68	144.40	164.86	3.48	98.46
1409	425.27	87.17	260.86	8.34	98.46
8815	664.61	1111.50	420.20	41.92	98.46
12130	90.83	34.97	39.25	3.30	98.42
8213	3583.64	1528.59	10080.02	1528.73	98.42
109	556.91	312.14	239.04	32.06	98.42
21637	28.01	19.25	-16.57	5.06	98.29
15819	42.77	22.12	-47.81	46.21	98.16
44	34.58	17.78	-3.41	3.04	98.16
6154	256.83	383.44	-1506.42	1100.74	98.12
13412	28.40	24.99	148.78	61.50	98.12
6720	68.84	1		21.68	98.12
5117		60.28	240.34		
5329	162.84 47.29	78.90 21.30	364.60 10.39	35.27 2.22	98.07 98.07
21866	109.05	73.32	379.42	172.12	98.03
14953	482.64		311.16	19.07	97.99
18350	90.05	76.56 49.29	301.03	46.70	97.99
2029	305.29	103.47	350.85	2.68	97.99
8837	358.84	91.24	173.81	19.49	97.95
25 7 21	83.34	54.43	240.21	44.54	97.95
16272	191.69	64.06	131.07	3.08	97.95
5969	1516.76	347.22	2916.15	354.10	97.90
1689	4338.65	2126.09	15982.97	5400.06	97.90
4232	137.19	62.23	27.42	11.32	97.90
3049	202.76	101.21			
18800	2711.70	1148.84	529.19 6417.82	69.46 556.95	97.86 97.86
14424	67.07	130.91	324.02	83.71	97.86
			477.43		
8849	222.06	71.45		62.77	97.77
25777	404.46	182.63 91.89	824.09	102.83	97.77
16902 23078	66.96		-172.67	27.28	97.77 97.77
	147.27	46.50	55.48	8.60	
5461	193.94	117.20	396.91	37.67	97.73
24814	171.74	33.51	92.18	13.93	97.73
10860	46.07	29.90 76.29	-6.09	2.93	97.69
1698	74.37		197.25	24.89	97.69
15408	193.87	58.16	69.92	11.05	97.69
17832	1948.88	851.66	5675.58	2032.21	97.65
7127	280.56	95.94	21.98	52.78	97.65
744	334.35	68.28	208.28	11.29	97.60
3081	387.30	84.71	227.05	31.49	97.56
18918	31.10	27.35	-32.80	10.09	97.56

TARIJE SP	CYCLOPHOSPH	////INB	AKK.	Dografia Na	44921-508977	
	: 6 hrs		Doc. No. 179332			
lemuliter			I noximean			
15154	267.93	62.29	131.15	21.56	97.52	
17771	768.98	305.15	1490.79	130.43	97.52	
20493	460.39	110.11	297.05	12.72	97.47	
24437	68.49	25.43	175.35	41.14	97.47	
8999	47.02	23.49	1.97	4.96	97.47	
15382	122.71	202.12	570.78	188.00	97.47	
8599	378.10	107.09	181.07	19.58	97.47	
17682	672.30	178.35	399.02	29.34	97.43	
22862	103.01	36.20	42.20	6.28	97.39	
20920	626.34	183.56	421.78	11.19	97.39	
17334	173.83	58.72	358.42	40.10	97.39	
4067	123.98	59.95	245.14	32.72	97.39	
17357	269.19	82.48	105.14	28.10	97.39	
23314	72.17	276.94	318.28	105.05	97.35	
19190	534.12	144.15	283.73	32.84	97.35	
16943	2759.04	872.63	5041.36	392.40	97.35	
16947	319.84	80.34	186.33	13.16	97.35	
6405	380.95	93.60	241.49	11.37	97.35	
9053	239.61	50.65	135.66	11.51	97.35	
25253	293.63	65.17	155.93	16.09	97.30	
1688	5256.94	3710.90	21624.32	7449.14	97.30	
19993	2319.53	566.29	3763.82	221.97	97.30	
20846	2377.64	663.94	3619.10	135.20	97.30	
22142	32.37	17.29	0.24	3.37	97.26	
17602	131.67	36.83	52.59	13.48	97.26	
18274	300.34	63.28	159.26	25.71	97.26	
15410	504.68	101.83	257.23	33.63	97.22	
7299	181.01	153.33	372.36	39.71	97.22	
6585	649.58	366.37	1390.92	91.78	97.17	
17426	537.84	84.20	386.94	14.48	97.13	
15190	1818.28	1175.28	4173.32	371.68	97.13	
13598	349.78	108.22	755.01	165.06	97.09	

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Timepoint(: DIFLUNISAL s): 24 lws			DOG: 1X	1921-5089W0 10. 1793397.1
ldendifier		NonToxSD	ToxMean 🐇	ToxSD	LDAScore :
15582	98.54	389.02	523.22	20.51	99.87
23699	325.81	88.09	984.09	67.68	99.79
1858	165.70	50.67	468.35	55.34	99.66
18687	415.65	168.60	1876.04	173.83	99.66
20810	1260.74	402.11	2154.96	30.08	99.44
23698	272.98	101.50	824.56	62.23	99.44
15906	73.43	105.60	328.56	63.09	99.40
21354	414.86	118.96	1273.64	189.63	99.36
16918	1181.89	410.28	2103.00	56.71	99.36
15048	861.35	213.04	1349.81	15.30	99.32
17758	126.22	53.45	567.25	114.93	99.14
23504	176.69	49.53	287.50	5.80	99.14
18686	468.60	214.77	2050.51	247.31	99.14
5351	711.19	145.06	979.89	15.97	99.06
18083	48.75	24.85	148.28	10.77	99.06
1977	142.19	40.33	293.78	26.17	98.97
20833	1254.24	350.22	1826.03	17.16	98.84
4012	650.38	267.26	1484.68	174.94	98.84
18250	1093.45	315.54	1684.15	30.30	98.80
26109	69.08	76.43	458.33	59.25	98.76
4049	23.28	65.36	63.64	6.15	98.76
23837	101.76	41.27	57.70	0.48	98.76
25679	901.31	261.02	1610.91	63.61	98.76
5887	79.37	88.10	285.80	15.29	98.63
23409	535.77	161.09	346.74	3.30	98.54
6380	135.92	123.74	459.26	82.61	98.54
2457	289.86	77.14	519.55	38.45	98.42
5667	744.60	177.74	1191.89	49.47	98.42
18293	770.57	235.09	1635.35	158.85	98.37
15579	32.24	115.84	189.18	29.09	98.33
18647	251.55	67.39	138.00	4.76	98.29
16849	114.09	44.40	66.30	1.36	98.20
15580	95.99	152.88	276.60	38.55	98.20
11205	594.93	193.66	371.24	5.25	98.20
17211	1440.25	553.34	2327.85	45.31	98.16
19244	1224.59	378.82	2305.53	135.95	98.16
1728	351.65	81.13	555.88	47.36	98.16
4010	965.06	407.94	2268.41	360.66	98.12
17563	1201.51	347.34	2039.09	80.83	98.07
19067	174.78	51.26	80.87	6.55	98.03
14763	38.55	133.68	500.64	63.55	98.03
17158	240.63	83.07	105.20	10.26	97.99
19727	1326.99	416.44	2052.69	41.43	97.90
2708	383.06	86.76	506.29	10.15	97.90
16204	755.43	187.45	1101.24	54.31	97.90
24748	-22.65	38.12	33.61	1.39	97.86
15239	567.21	132.24	803.19	32.26	97.82
22052	256.34	73.77	451.80	35.28	97.77
20715	135.54	60.66	373.16	45.27	97.73
19268	940.57	273.83	1595.25	83.86	97.73
17686	1016.62	266.11	1780.84	121.15	97.69
20986	36.18	25.28	113.63	23.08	97.65
3027	1127.48	331.63	1758.65	53.48	97.65
23849	287.19	137.85	391.28	6.74	97.60
4952	108.82	48.80	171.37	6.32	97.60

-414-								
TARIE G	e: Diflunisal	in the same of		alta Ma	1921-5089WO			
	((s): 24 hrs		معروب		1798897.1			
ldendifter	NonToxMean		TorMean		LDAScore			
1814	171.85	47.69	123.88	1.67	97.56			
20839	1043.73	290.47	1565.67	42.99	97.52			
16190	288.78	81.68	502.75	35.87	97.52			
15875	1183.10	392.02	1973.54	86.87	97.35			
16701	830.81	197.66	1528.91	267.69	97.35			
15106	1887.41	709.31	3311.08	123.87	97.26			
3434	326.00	140.86	131.21	15.66	97.26			
21729	582.55	246.48	1512.04	499.79	97.26			
19952	67.28	24.42	23.96	3.74	97.26			
20818	681.56	422.22	1663.94	317.99	97.17			
20149	1324.01	582.48	2795.60	215.42	97.17			
14959	619.74	147.94	1010.98	113.74	97.13			
16148	762.99	195.12	1313.23	188.91	97.13			
24886	1263.69	371.87	2046.76	132.13	97.13			
10878	952.87	253.83	1373.59	38.24	97.09			
8946	207.89	87.83	74.88	7.06	97.09			
19477	112.98	54.74	250.89	20.66	97.00			
15468	734.22	179.64	1032.19	49.88	97.00			
10109	1045.52	288.55	1597.93	74.73	96.96			
3924	182.19	68.67	273.97	5.41	96.92			
18918	31.02	27.50	-11.16	3.27	96.88			
17729	878.67	224.12	1508.83	163.08	96.83			
14695	1736.29	632.39	2885.22	195.16	96.79			
9799	145.87	50.77	85.38	3.80	96.79			
20925	337.86	103.37	649.53	101.58	96.79			
25501	102.21	56.92	20.07	8.06	96.79			
7062	686.14	173.76	1144.72	115.88				
5398	0.40	12.35	33.62	24.14	96.79 96.75			
20711	43.78	42.18						
373	34.25	61.71	150.73	22.20	96.75			
16929			167.08	44.08	96.70			
20817	956.64 1073.65	234.98 677.30	1405.43 2215.68	58.88 400.07	96.70			
4291		I			96.70			
4291 23336	295.13	101.25	151.24	8.90	96.70			
	141.91	43.60	268.14	38.59	96.66			
23270	209.26	61.36	302.15	9.89	96.66			
4259	700.25	158.48	1007.49	53.89	96.66			
18509 1694	418.06	82.28	584.02	27.38	96.62			
	1145.19	337.82	1796.01	81.50	96.58			
4011	459.62	198.95	896.31	148.45	96.53			
6949	1.61	27.50	36.30	10.98	96.49			
4713	107.84	37.11	71.04	3.04	96.40			

TABLE 5R: HYDRALAZINE Timepoint(s): 6 hrs			Doc. No. 1798397.1		
ldentifier	MonToxMean	NonToxSD .	ToxMean	ToxSD	LDAScore
23230	381.93	101.69	164.86	3.08	99.83
7299	180.37	151.55	684.70	26.37	99.66
11005	71.62	20.88	31.26	0.53	99.66
18715	190.60	53.82	312.84	2.17	99.66
18713	300.35	70.22	564.29	26.19	99.62
9306	45.76	18.52	121.89	7.27	99.62
19004	788.98	306.09	794.75	1.65	99.57
9525	5.08	44.31	64.20	2.12	99.57
19712	98.45	32.90	58.61	0.21	99.53
16203	66.52	16.97	45.27	0.17	99.53
20513	58.53	27.11	242.01	35.67	99.49
5918	26.70	40.82	72.84	4.20	99.44
14479	473.36	112.67	261.32	4.86	99.44
4194	34.33	17.04	74.46	0.76	99.40
8948	217.22	104.91	343.57	1.20	99.40
15015	503.71	91.72	339.92	3.82	99.36
22746	534.33	197.34	312.42	2.15	99.36
13235	16.43	13.42	75.50	8.64	99.32
90	93.87	50.03	147.79	0.44	99.32
16069	59.01	26.48	31.64	0.25	99.32
20523	549.70	152.39	1021.93	29.01	99.23
15872	155.66	71.91	53.38	1.82	99.23
3513	171.87	35.63	273.79	6.58	99.19
7452	8.04	13.04	37.85	2.68	99.19
15059	92.43	24.57	82.57	0.16	99.14
8597	241.04	57.82	387.68	12.70	99.14
20849	277.59	87.41	309.60	0.68	99.06
13298	101.48	29.62	51.68	1.68	99.06
11406	242.13	50.08	177.73	0.85	99.02
7071	133.86	64.50	98.99	0.88	99.02
23189	330.92	72.72	460.60	3.57	98.97
15411	303.87	90.07	657.96	59.65	98.97
8692	1045.40	309.78	638.87	9.02	98.97
22614	17.91	87.18	86.87	14.83	98.93
4969	-18.84	34.82	77.79	30.12	98.89
3081	387.16	84.82	227.51	15.13	98.89
15231	100.29	44.00	233.89	22.19	98.85
15942	230.65	65.79	294.11	1.11	98.85
18406	33.60	18.22	70.49	1.79	98.85
16024	239.14	63.32	487.33	52.09	98.85
2539	46.96	51.90	304.80	46.42	98.85
26119	124.62	46.64	176.21	1.26	98.80
6723	200.38	72.05	91.78	10.99	98.76
21878	144.05	35.94	106.39	0.58	98.76
8664	106.71	250.43	160.02	4.54	98.72
21014	155.47	80.73	369.94	37.53	98.72
14842	122.80	45.94	280.85	20.55	98.72
13093	700.37	165.48	367.14	36.76	98.72
20404	53.45	44.01	336.92	98.70	98.72
15126	792.34	224.59	1256.68	25.48	98.67
4948	153.55	61.38	307.30	11.57	98.63
6844	123.76	58.60	32.99	3.47	98.63
16025	150.51	42.20	268.08	8.85	98.59
7615	90.87	40.61	221.25	17.10	98.55
22575	20.61	15.66	10.19	0.32	98.50

	TABLE 57: HYDRALAZINE AXY. Docket No. 44921-5030						
	((s)): 6 hrs				1798897.1		
ldentifier	NonToxMean	Montoxed	ToxMean	ToxSD	LDAScore :		
15636	39.88	26.09	4.00	1.07	98.50		
15885	93.27	28.63	182.42	15.91	98.46		
5355	1164.75	345.23	1050.46	4.54	98.46		
13151	711.00	343.67	1290.08	71.04	98.46		
19195	1555.05	374.36	1121.86	8.23	98.42		
6606	249.66	121.42	743.36	159.90	98.42		
2888	2015.72	588.67	1201.76	29.67	98.42		
20405	34.08	29.57	195.15	62.34	98.42		
7197	197.58	84.65	309.34	7.55	98.42		
1215	67.05	49.57	246.66	40.98	98.42		
17479	157.69	34.71	89.24	4.57	98.37		
22733	21.90	14.11	64.22	8.36	98.37		
1920	426.50	112.57	740.10	44.20	98.37		
8745	57.56	19.95	100.05	2.83	98.33		
22915	171.30	47.15	114.08	2.66	98.33		
25587	24.39	16.37	17.05	0.15	98.33		
13259	68.80	25.82	161.38	28.29	98.33		
17468	396.41	77.87	278.70	4.37	98.29		
14405	465.44	308.54	1276.71	188.93	98.25		
14861	48.74	17.37	75.52	1.15	98.25		
3027	1129.44	333.52	1205.87	5.52	98.25		
1214	165.46	52.58	424.14	92.45	98.20		
11158	1023.54	302.35	1115.38	6.37	98.20		
20202	624.37	196.92	959.56	18.31	98.20		
18290	275.55	78.31	420.92	7.75	98.16		
21527	239.85	60.68	369.29	12.50	98.16		
24885	1107.60	334.53	1092.86	5.80	98.16		
23689	0.64	11.77	22.99	3.94	98.16		
8869	8.35	21.59	31.83	0.60	98.16		
17502	147.76	53.99	264.34	13.12	98.16		
13203	-15.66	28.87	51.81	11.94	98.12		
25971	106.47	34.61	50.27	2.15	98.12		
13095	60.49	15.80	32.26	1.41	98.12		
26036	34.25	17.22	19.79	0.49	98.12		
17570	230.73	71.54	273.93	2.11	98.08		
22543	689.99	222.20	447.42	9.98	98.08		
17312	26.98	34.41	42.60	0.40	98.08		
16026	221.52	75.94	411.54	26.17	98.08		
5684	333.75	76.46	482.85	10.21	98.03		
15879	374.14	93.39	276.43	3.61	98.03		

TABLE 58: Timepoin((s	IFOSFAMIDE 5): 6, 24, 43, 144 h	is a		<u> </u>	1921-5039WC 10. 1793397.1
ldentifier 🐃	NonToxMean **	NonTox&D	ToxiMean	ToxSD	LDAScore
19252	647.94	142.98	520.93	34.13	89.74
1622	2071.83	907.04	1625.21	143.87	88.88
16148	767.65	197.63	575.47	57.95	86.32
17779	1952.57	790.85	1617.53	137.53	85.80
12932	153.85	49.70	108.04	13.75	85.63
11158	1027.66	302.42	759.65	56.80	85.37
10947	1800.32	754.28	1416.69	180.51	85.28
19408	2011.57	680.23	1580.63	160.66	85.24
16013	58.91	18.74	71.23	5.22	85.11
19254	241.32	76.35	239.54	14.43	85.06
17886	1504.90	495.37	1122.51	90.54	84.72
16895	2027.29	906.56	1603.00	189.71	84.63
18300	450.17	150.80	316.95	39.08	84.33
8211	2862.98	1618.85	2138.69	273.26	84.29
23710	1141.57	367.03	897.82	79.04	84.20
11954	3141.99	1699.73	2471.68	314.40	83.90
1853	2014.76	833.49	1614.19	220.80	83.90
14695	1746.72	637.07	1297.75	126.83	83.77
8212	2267.04	1143.40	1689.07	186.62	83.42
13976	453.36	377.22	783.29	266.52	83.38
14997	2596.32	1029.67	2216.17	249.47	83.20
23709	2500.59	1207.33	1994.07	261.44	83.16
22592	234.68	168.43	397.15	108.13	83.08
18142	2001.91	840.95	1479.52	180.12	82.99
9135	719.00	131.99	601.54	43.95	82.90
14694	2326.49	1072.07	1849.47	243.07	82.69
18810	1189.60	320.47	979.08	72.74	82.60
18077	2627.05	1191.54	1778.68	232.31	82.56
15125	1303.70	426.26	1007.41	95.34	82.55
20751	706.77	161.80	835.79	96.53	82.51
44	34.24	17.91	48.60	8.82	82.47
23544	1515.02	495.50	1325.70	110.75	82.17
19993	2320.17	574.10	2616.44	234.66	82.03
9942	443.45	98.80	518.47	38.50	81.95
17682	673.25	179.30	543.60	51.66	81.95
23574	2282.20	973.76	1808.58	200.84	81.82
6815	1138.28	344.89	837.60	73.20	81.65
11050	671.41	139.50	791.75	84.37	81.60
1247	1310.50	504.69	915.02	93.84	81.60
18078	1152.53	545.02	1014.06	139.87	81.56
1801	97.60	29.90	123.54	20.25	81.43
12901	1613.41	414.49	2126.00	326.75	81.39
20035	182.08	110.21	229.47	33.04	81.39
17118	53.61	19.28	66.07	6.68	81.30
17204	1733.32	660.21	1299.09	155.20	81.26
23847	47.46	36.72	90.72	17.29	81.21
10500	24.88	31.73	53.21	18.49	81.17
8347	76.96	46.62	110.08	23.49	81.13
3015	2318.93	997.42	1869.25	225.78	81.04
20832	722.84	197.13	591.89	72.43	81.00
18615	521.90	148.77	363.34	55.31	80.95
5989	269.37	71.43	354.20	68.39	80.91
19894	40.81	20.31	62.51	16.62	80.83
18076	2548.69	1193.37	1768.87	244.55	80.83
21423	1355.89	429.81	1080.04	115.43	80.78

	S: IFOSFAMIDE M(s)): 6, 24, 48, 14		AMy. D		4921-5089WC No. 1798897.1
	MonToxMean		areaMissoriii		
1523	89.47	23.47	100.14	7.66	80.70
11991	57.24	22.13	76.61	11.20	80.65
23109	2089.70	930.42	1566.07	197.88	80.44
19727	1331.92	420.23	1162.29	117.79	80.44
18451	1434.72	418.40	1801.12	288.28	80.39
23884	41.09	33.35	54.46	12.98	80.39
472	662.53	178.22	500.21	63.31	80.35
11153	1474.65	500.35	1320.86	134.33	80.35
23125	4577.67	2547.90	5632.15	1198.40	80.31
12598	545.25	95.85	644.01	73.00	80.31
915	30.40	17.20	53.68	14.78	80.26
6808	698.18	193.62	893.93	94.12	80.26
10260	85.57	33.38	94.71	10.64	80.18
25545	94.20	49.43	150.75	32.71	80.09
23660	1281,20	383.06	1072.18	89.30	80.09
15410	504.96	102.87	427.16	47.59	80.05
977	16.73	10.91	41.99	18.19	80.01
15137	1520.02	496.49	1254.30	128.20	80.00
26109	69.41	79.57	138.16	53.60	79.96
11136	1003.50	311.85	725.42	110.05	79.92
4217	519.77	126.27	592.35	45.95	79.92
13480	650.24	137.90	533.92	69.48	79.83
15535	448.65	83.59	358.13	46.40	79.79
15426	411.88	86.42	342.56	30.90	79.66
16012	73.15	32.18	78.40	10.56	79.65
4849	773.73	168.65	929.55	91.93	79.61
17765	1296.75	460.31	914.95	107.23	79.61
23967	383.11	92.35	484.27	65.27	79.53
9905	673.59	140.94	588.24	39.94	79.53
1583	30.08	18.99	43.08	7.71	79.53
1743	29.18	16.19	50.78	15.93	79.40
15446	370.71	93.07	266.06	61.63	79.40
18905	1363.50	302.26	1680.54	182.30	79.36
24049	1519.32	446.29	1776.05	182.78	79.31
24626	1504.58	431.27	1251.16	114.87	79.31
820	2467.06	1167.88	1920.44	339.35	79.22
1684	2833.17	1621.88	1212.60	870.83	79.22
21373	373.62	85.16	332.82	32.95	79.22
16211	2261.75	1141.46	1652.93	320.55	79.18
16521	285.41	90.61	359.56	55.31	79.14
22661	1309.12	421.45	1003.73	117.38	79.01

Table 5T: Indomethacin Timepoint(s): 43, 72 hrs			Aity. Docket No. 44921-5039WC		
Timepoint(s)): 45, 7/2 hrs ::		 		<u> </u>
	NonToxMean -		24.	MANAGEMENT BEAUTY OF THE PARTY	
155	21.42	16.98	101.88	13.61	99.53
154	112.29	36.99	249.23	25.25	99.27
16173	14.63	13.08	179.81	66.16	99.18
13614	340.21	88.42	786.72	118.82	99.01
1850	46.85	347.25	309.01	159.03	98.93
22499	8.20	11.83	58.62	9.91	98.88
1893	29.44	20.17	163.67	70.52	98.84
1221	0.23	15.64	193.08	98.35	98.80
21445	0.99	14.21	203.14	86.78	98.75
1854	43.31	289.94	282.61	143.16	98.67
25517	38.27	31.10	173.50	50.20	98.58
19710	43.43	20.81	132.45	42.06	98.54
6431	51.07	32.99	209.15	53.95	98.50
2457	288.90	75.46	543.74	54.46	98.45
7299	177.44	143.66	797.13	276.35	98.41
23964	12.50	18.01	63.92	17.10	98.37
1943	31.78	14.21	86.66	15.03	98.15
13615	253.23	68.59	560.03	77.71	98.15
20713	215.61	108.00	606.67	98.17	98.11
24237	56.39	37.15	281.98	103.56	98.07
8565	31.85	16.10	105.94	38.19	98.02
7540	154.84	89.44	535.02	179.10	98.02
1845	-7.69	24.61	109.59	48.06	98.02
18684	137.77	49.02	305.11	48.29	97.98
7858	-4.71	7.47	47.48	40.75	97.98
15408	192.19	56.59	376.55	44.39	97.98
10281	172.58	185.66	567.51	296.37	97.94
18867	103.82	49.82	263.24	51.22	97.94
18353	112.33	68.84	349.21	48.50	97.94
20715	134.85	59.63	354.19	34.91	97.90
6551	576.57	201.46	1150.85	135.93	97.90
7665	282.66	95.03	609.80	128.48	97.90
20868	22.06	17.54	98.66	30.62	97.85
343	28.72	32.40	151.55	21.38	97.85
20869	21.84	21.14	111.52	32.81	97.85
20711	43.08	40.65	197.85	41.42	97.77
16521	283.95	85.02	655.14	127.79	97.59
21444	19.93	43.02	180.28	49.29	97.51
21683	33.96	22.93	127.35	39.46	97.47
3180	309.65	80.63	524.23	45.30	97.47
1942	17.90	24.48	99.74	38.19	97.47
14184	113.17	49.05	251.46	48.41	97.42
1894	202.61	73.39	421.42	64.64	97.42
15851	200.88	151.27	525.10	94.05	97.38
20700	85.68	416.32	391.89	193.02	97.38
1 74 9	234.65	200.18	431.09	62.04	97.38
5094	138.21	54.26	356.45	88.21	97.38
2555	96.41	46.12	235.98	52.95	97.34
3260	193.81	71.46	443.67	108.75	97.34
19012	483.92	149.86		98.29	97.34
5887	77.25	81.17	930.52		
20041			491.60	169.72	97.16
16007	192.79	82.66	490.91	119.03	97.16
	26.63	17.31	92.50	26.48	97.04
21653	236.04	61.07	399.34	79.22	97.04
13004	142.59	42.21	275.32	36.65	96.99

TABLE ST:	INDOWETHAC (s): 48, 72 lws	N I	Ally. Docket No. 44921-5089W Doc. No. 1798397		
lejanjijar Tumasanin	NonToxMean				
19387	667.36	146.95	1017.20	82.56	96.99
2554	53.34	18.18	111.59	20.52	96.99
4661	307.32	90.25	543.97	56.72	96.91
21467	13.64	131.26	140.69	60.68	96.86
18352	162.12	85.33	437.38	90.35	96.86
15191	2030.38	1211.92	434.31	67.70	96.86
24183	68.30	35.60	194.82	48.12	96.82
4748	110.39	127.49	266.50	52.38	96.82
19711	83.58	22.63	149.45	21.04	96.82
848	18.09	11.05	45.15	4.76	96.65
10015	231.63	76.39	422.68	89.46	96.65
11708	319.08	90.36	587.03	103.59	96.65
22321	101.90	68.06	234.02	51.42	96.61
1597	50.19	41.89	187.05	63.28	96.61
6120	511.17	159.69	914.06	106.73	96.52
20714	178.33	78.97	430.20	87.05	96.52
24200	421.16	141.29	847.55	177.88	96.52
3316	8.48	11.20	32.02	7.24	96.48
14595	87.99	36.42	175.31	16.47	96.39
21654	362.19	114.34	536.63	59.58	96.22
22479	433.07	146.06	846.76	137.55	96.18
18687	418.98	188.38	661.54	59.97	96.13
5572	330.93	144.21	692.30	98.63	96.09
3020	292.14	93.53	549.45	92.30	96.09
25366	52.69	46.59	140.27	34.35	96.01
11183	98.16	39.08	183.02	34.74	96.01
19145	363.77	71.33	513.30	38.33	95.96
15409	399.23	98.94	637.09	62.27	95.88
17950	64.06	23.04	112.35	10.61	95.79
16945	934.12	162.35	1222.61	66.01	95.79
16917	755.96	258.94	1351.72	135.35	95.75
9286	205.05	59.59	354.73	51.76	95.66
18217	15.31	19.01	63.19	16.35	95.62
14185	203.46	94.18	411.57	81.62	95.62
16646	28.33	20.91	71.67	15.31	95.58
23837	100.89	40.39	206.26	37.38	95.53
21066	43.85	16.28	90.93	15.15	95.53
18068	82.30	21.15	125.33	8.28	95.49
21410	183.82	61.54	326.34	61.71	95.45
18355	56.83	27.50	129.21	39.86	95.36
6044	327.15	91.24	520.74	70.39	95.36
23145	44.14	20.10	96.23	26.84	95.32
16859	122.48	51.26	241.14	54.89	95.27
15246	63.40	20.73	113.00	17.39	95.19
24019	27.09	25.66	86.45	18.38	95.19

	FILTHINIM CHROL		Ally. Docket No. 44921-5039W			
	<u>s): 120 brs</u>				No. 1798897	
dentifier 🦠	MonToxMean =	NonTox&D 🐇	ToxMean	ToxSD	LDAScore	
24019	27.07	25.30	127.00	8.27	99.61	
4421	418.40	87.55	639.64	12.93	99.57	
13641	47.49	30.07	145.93	13.26	99.53	
463	619.17	296.16	1455.45	46.93	99.53	
6300	61.51	20.88	147.76	15.29	99.49	
13974	296.90	139.79	1260.41	77.75	99.49	
3244	127.45	30.43	40.35	2.64	99.36	
10464	131.43	35.68	41.70	2.59	99.36	
1462	407.93	201.27	1216.38	60.80	99.36	
8525	246.32	66.03	97.58	6.10	99.27	
4227	72.35	65.02	265.04	34.60	99.23	
25741	185.55	57.43	490.81	62.92	99.23	
1224	3.70	11.53	50.50	8.69	99.23	
3880	679.69	204.44	295.39	24.01	99.14	
14261	39.73	17.42	113.90	16.40	99.14	
18472	1551.82	378.09	827.52	66.71	99.10	
15363	446.56	133.55	187.16	15.68	99.06	
3598	139.25	41.61	362.49	73.57	99.01	
5176	217.77	121.65	603.36	60.94	98.93	
14754	99.93	24.11	228.47	44.23	98.93	
20849	276.44	85.41	565.38	36.71	98.93	
15955	750.17	198.73	371.84	30.56	98.89	
14633	97.72	88.30	-1.97	5.36	98.84	
15371	378.04	55.30	593.66	40.88	98.80	
17342	202.36	532.80	480.79	60.48	98.80	
20809	317.42	60.81	558.00	56.79	98.80	
16650	256.32	82.88	614.21	64.30	98.80	
18109	19.25	21.24	92.33	9.99	98.76	
18286	10.51	10.22	41.67	5.33	98.76	
24049	1526.25	442.84	773.94	52.98	98.76	
16245	386.32	115.78	64.71	31.63	98.76	
23651	656.57	627.85	1600.29	142.54	98.76	
574	335.41	164.90	679.82	42.41	98.71	
20099	81.48	29.97	171.81	18.17	98.71	
1976	24.82	21.34	269.98	118.48	98.71	
23294	244.42	55.35	115.25	8.35	98.71	
3733	340.43	153.36	790.69	47.48	98.67	
23957	68.22	41.24	264.04	43.03	98.67	
3348	496.09	130.59	245.12	25.78	98.67	
20697	1418.84	278.90	908.67	36.86	98.67	
260	415.71	110.33	903.01	137.68	98.63	
9321	313.57	55.98	464.14	23.58	98.63	
3597	240.21	54.94	523.47	111.57	98.63	
7159	699.06	218.26	1124.72	53.18	98.63	
8103	58.33	28.29	149.64	17.07	98.59	
698	1169.95	220.06	1885.00	141.61	98.59	
24648	40.24	20.33	6.76	1.40	98.59	
4020	277.14	68.14	152.09	12.77	98.54	
5246	63.64	21.09	87.26	0.84	98.54	
4115	68.51	36.48	345.37	126.46	98.54	
21997	28.94	25.26	95.22	19.06	98.54	
960	191.35	46.71	93.10	7.12	98.50	
2321	100.55	58.87	629.47	166.32	98.50	
20886	711.00	279.64	2114.90	382.05	98.50	
6354	280.15	119.74	-118.26	64.76	98.50	

	-422-								
TABLE 5U: LITHIUM CHLORIDE AMy. Docket No. 44921-5039WO Timepoin((s)): 120 hrs Doc. No. 1793397.1									
		NonTox SD	ToxMean						
12376	19.26	124.98	83.39	19.10	98.50				
17106	81.66	24.82	36.82	2.74	98.50				
18473	2589.69	916.99	1318.66	52.24	98.50				
9254	246.49	48.78	393.24	20.14	98.46				
1340	191.26	49.84	289.47	26.83	98.46				
17162	6.69	16.97	58.11	15.91	98.46				
6806	1233.36	345.85	586.60	42.84	98.41				
25840	-2.65	12.34	54.97	27.90	98.41				
20887	784.36	296.51	2396.08	485.51	98.37				
1372	199.45	51.98	391.46	35.86	98.33				
18349	140.08	38.43	262.92	23.80	98.33				
14989	500.36	106.29	910.22	99.63	98.33				
4049	23.03	65.14	113.93	24.17	98.29				
11483	65.04	45.49	132.25	10.90	98.29				
9867	29.07	22.58	-38.05	11.94	98.29				
13411	789.51	324.70	239.08	27.46	98.29				
25709	267.20	83.61	591.14	81.21	98.29				
17160	1038.70	287.67	1872.62	168.15	98.24				
12673	37.66	22.70	116.03	13.02	98.24				
25306	27.76	21.91	-55.70	19.66	98.24				
16349	47.04	14.11	94.68	13.74	98.24				
13392	190.93	48.80	355.59	34.39	98.24				
4048	-6.11	32.04	34.41	10.44	98.20				
2915	87.49	36.50	176.88	18.22	98.20				
15761	90.06	42.30	20.08	3.24	98.20				
111	590.89	237.37	1115.91	164.07	98.20				
24695	63.72	21.72	13.53	4.47	98.16				
19152	164.93	52.20	401.36	74.04	98.16				
24597	585.04	105.40	935.83	83.53	98.16				
21527	239.33	59.62	438.09	50.65	98.16				
3776	377.17	93.88	689.18	68.23	98.16				
5626	26.84	18.16	84.22	9.81	98.11				
3075	524.14	179.66	209.94	19.92	98.11				
15511	214.84	128.36	897.69	308.94	98.11				
6386	20.45	18.01	-5.75	1.87	98.07				
134	69.64	33.24	7.05	4.13	98.07				
19665	91.84	47.87	296.74	33.57	98.07				
10984	2015.62	568.01	964.72	141.17	98.03				
15247	637.80	190.94	1369.01	148.83	98.03				
2905	246.54	108.85	410.67	11.79	98.03				
13343	210.51	55.05	72.01	21.01	98.03				

TABLE 5V: (I	MERCURIC CHLO	RIDE Alty. Docket No. 44921-5 Doc. No. 179			
<u>timagema(e)</u> Identifier	NonToxMean		ToxMean	ToxSD	LDAScore
8665	332.39	171.86	5197.14	2943.02	99.44
1475	136.94	103.32	3940.87	1974.36	99.18
20035	178.08	98.83	635.50	155.87	98.10
19723	74.87	55.61	273.03	109.44	98.02
15191	1988.23	1177.92	5041.11	935.19	96.55
14139	80.54	27.01	25.59	13.59	95.47
8664	86.42	70.91	2082.06	1352.81	95.23
12331	594.25	155.97	254.07	138.26	94.96
17734	119.63	75.92	566.88	280.56	94.84
23579	734.95	158.66	430.76	80.03	94.78
23983	544.27	150.95	220.82	131.74	94.48
14138	84.72	29.10	33.29	12.42	94.31
16518	826.45	273.09	2395.53	930.51	94.24
17635	651.30	148.78	358.62	92.32	93.79
7196	178.22	77.07	494.09	205.98	93.68
15850	1203.52	323.32	2507.97	545.30	93.68
24235	433.60	163.10	1038.60	349.80	93.55
21445	2.08	22.76	30.67	12.91	93.51
24649	122.93	28.55	76.77	12.19	93.49
5867	166.42	48.13	259.79	50.17	93.49
3348	497.46	129.58	259.25	78.88	93.45
15848	1313.68	402.72	2886.50	676.70	93.42
19768	706.30	166.91	1328.43	290.35	93.42
17161	1144.51	381.55	2737.15	723.56	93.33
1004	108.90	32.28	56.68	10.23	92.84
15190	1789.26	1124.32	5409.42	1117.57	92.73
14595	89.23	36.64	26.57	16.82	92.67
15189	1722.10	1120.12	5391.29	1462.09	92.60
15300	135.69	130.01	397.39	125.33	92.60
23314	67.33	271.41	622.50	259.52	92.43
	37.59	67.32	147.90	49.32	92.39
15301 18944	202.65	68.48	458.00	181.57	92.30
	27.38	53.50	71.68	29.34	92.26
6054	209.02	74.04	85.92	29.90	92.24
13642		96.00	695.07	177.40	92.17
23230 17211	378.13 1433.79	549.05	2360.95	279.33	91.85
3493	58.50	20.07	107.74	26.19	91.82
	283.86	49.04	166.00	50.35	91.78
23825			95.46	32.77	91.74
18564	219.22	49.05 81.92	233.52	34.79	91.68
11680	360.19		2585.84	579.12	91.65
17765	1277.82	438.77	380.00	223.17	91.57
16982	138.22	290.89	196.18	80.11	91.26
13610	357.65	69.18		18.36	91.26
21993	76.47	25.96	129.83	106.17	91.01
8927	745.07	166.28	377.77		90.91
11871	55.24	129.01	197.33	57.60	90.88
11050	670.06	136.16	972.43	147.61	90.88
13507	422.59	99.12	659.30	138.95 25.82	90.83
9271	102.44	54.22	21.68	_ I	
19031	59.98	47.58	145.41	39.97	90.79
24577	1168.59	364.98	2135.75	393.09	90.79
10182	2.45	99.29	78.59	59.67	90.75
18300	451.47	147.74	136.58	90.13	90.75
18259	207.76	78.23	1353.31	770.76	90.72
1928	485.55	106.87	208.21	110.11	90.66

TARIESW.	MERCURIC CHLO	aine i i	alan No. A	14921-5039WO	
illimepoint(s)	ያ 35		Doc. No. 1793397.1		
ldendifer	NonToxMean	NonToxSD	ToxMean	ToxSD	LDAScore
6632	171.87	58.89	288.22	64.66	90.53
13611	273.95	107.44	57.12	66.20	90.45
25098	43.26	30.01	276.68	179.66	90.37
22539	71.67	46.78	-27.28	37.71	90.36
20945	897.15	248.57	1469.22	277.42	90.32
19678	110.11	57.05	-19.38	53.40	90.32
23567	52.11	60.74	235.62	105.54	90.29
23868	159.82	189.43	1997.17	1212.75	90.24
812	157.29	34.98	83.70	36.70	90.19
23872	44.34	60.45	587.85	460.34	90.16
21372	276.72	65.43	134.32	44.54	90.10
18611	1409.94	446.15	2438.75	459.58	90.10
21306	126.88	54.43	52.78	18.76	90.09
3808	166.04	48.80	407.80	212.39	90.07
12031	145.86	38.78	225.16	30.38	90.06
23869	36.50	52.75	579.45	394.24	90.03
3015	2289.41	964.34	4534.92	1150.81	89.97
17908	60.79	49.95	325.35	238.47	89.90
25539	125.13	35.09	45.76	23.16	89.89
3473	120.74	34.02	69.52	20.97	89.83
2536	397.05	123.98	201.50	73.82	89.83
23826	344.12	55.78	225.14	47.99	89.80
9114	870.77	222.15	416.13	221.53	89.80
1639	97.37	22.99	62.34	12.37	89.70
20920	617.66	155.43	1397.16	623.36	89.60
20350	156.72	49.50	53.00	45.87	89.58
6615	279.24	84.15	96.62	96.14	89.50
19952	67.58	24.21	24.03	11.92	89.41
8237	102.59	36.73	179.81	51.98	89.37
11841	886.79	192.09	1632.36	479.53	89.34
2310	-47.82	40.85	36.34	37.62	89.32
15796	337.30	87.85	156.28	76.95	89.24
22681	216.80	177.12	918.02	459.53	89.21
22543	694.25	216.61	217.39	263.73	89.20
19433	110.29	59.85	445.41	230.04	89.17
22540	1828.95	538.43	794.41	586.09	89.07
17473	421.95	90.69	764.49	230.12	89.04
15875	1177.56	384.90	1982.41	472.79	88.98
18396	74.08	26.23	124.23	18.90	88.94
19	336.88	73.75	628.54	175.62	88.82
25567	456.18	167.99	922.98	373.29	88.65
20728	538.10	99.95	327.92	109.45	88.59
24351	-2.24	8.36	22.30	16.23	88.56
9053	240.22	50.14	146.06	39.78	88.55
18305	1364.98	457.67	2297.19	462.10	88.51

TABLE 500	RAMIDRONAT	Ē	- Ally. Doc	ket No. 44921-5089000	
Timepoint	(s)): 24 livrs				lo. 1798397.
dendifter !	NonToxMean	NonToxSD	ToxMean	TOXSD #	LDAScore
439	55.65	23.09	35.13	0.15	99.49
381	23.05	20.96	4.85	0.20	98.80
1439	233.12	46.90	167.13	1.64	98.63
24501	526.65	117.76	477.38	1.09	98.50
815	1161.46	344.32	1840.91	46.72	98.50
21723	24.61	15.80	18.81	0.26	98.37
25907	19.98	25.22	27.99	0.55	98.33
1440	320.37	110.50	249.77	1.69	98.33
3950	91.09	33.83	64.54	0.45	98.29
1145	59.20	26.43	36.98	0.67	98.25
20257	130.93	45.11	78.91	1.23	98.20
12781	189.97	49.11	273.46	3.66	98.12
20427	796.69	192.75	1119.78	17.38	97.90
16938	1322.80	411.98	1887.43	31.46	97.90
1324	63.66	31.18	28.89	2.88	97.82
16584	115.91	45.04	161.39	2.19	97.73
17102	47.91	18.57	30.16	0.64	97.73
10227	447.34	108.01	244.96	13.58	97.69
1310	152.06	33.32	86.35	7.14	97.69
8476	3812.40	1012.46	4862.16	57.22	97.65
22051	151.18	44.26	223.47	4.06	97.65
5049	297.65	66.21	180.38	7.11	97.60
16323	68.35	35.92	36.63	1.47	97.56
6654	166.71	45.74	158.83	0.92	97.43
1651	878.67	240.75	692.24	5.18	97.39
16192	41.41	14.72	34.35	0.38	97.31
12343	50.68	18.55	31.84	0.83	97.31
691	133.51	42.32	62.52	3.81	97.26
17635	648.73	151.15	484.36	6.12	97.22
9286	205.78	60.59	319.19	10.41	97.22
14800	36.39	23.63	34.69	0.41	97.22
23888	105.90	50.58	91.90	1.08	97.13
5969	1521.63	357.02	1485.82	9.84	97.13
3475	384.13	98.77	465.98	4.26	97.05
11174	54.15	51.85	121.54	7.12	97.01
26119	124.47	45.67	234.96	195.06	97.01
18250	1094.73	316.96	1382.93	14.86	96.96
25069	133.97	60.97	77.32	2.29	96.88
18135	133.61	28.59	96.51	1.97	96.88
21742	33.31	18.06	25.78	0.33	96.83
25702	579.55	128.85	805.98	44.29	96.79
10936	207.62	43.60	191.02	1.12	96.79
12342	96.27	49.48	87.13	1.04	96.75
19976	47.03	15.83	30.77	0.70	96.75
9620	530.95	116.93	770.14	47.74	96.71
20810	1262.30	404.25	1842.66	50.26	96.71
24721	203.81	53.70	187.98	1.38	96.71
14967	49.34	21.65	19.69	1.41	96.58
1309	43.01	16.54	30.28	0.59	96.54
730	112.34	33.32	65.78	2.57	96.49
15876	1174.15	310.95	1565.93	37.98	96.49
16482	195.58	40.06	132.17	5.76	96.45
5654	41.42	24.77	86.94	5.20	96.45
15850	1216.75	351.89	1264.63	16.42	96.45
4259	700.70	159.00	935.48	29.71	96.36

TABLE 5W	7: PAMIDRONAT	re Co	Ally, Do	eket No. 4	4921-5039\\\	
	(s): 24 hrs		Dock No. 1798397.			
ldentifier	NonToxMean :		ToxMean	ToxsD	LDAScore	
7010	309.13	55.63	255.94	3.25	96.36	
18880	36.99	12.90	31.14	0.58	96.32	
3007	126.64	42.37	192.91	7.32	96.28	
1375	84.74	20.63	55.45	2.98	96.28	
19244	1227.03	383.14	1715.96	64.01	96.28	
6595	76.76	35.41	102.22	2.83	96.19	
18126	701.66	166.62	548.76	14.64	96.19	
10869	11.30	60.29	123.29	15.83	96.15	
15239	567.46	132.45	783.53	43.73	96.11	
4241	114.51	43.66	203.36	162.90	96.11	
12360	55.02	27.32	35.03	1.44	96.11	
11687	25.88	23.50	35.95	1.69	96.11	
5492	97.03	47.49	24.35	5.81	96.11	
9671	124.41	52.83	135.25	1.91	96.11	
9410	62.57	24.27	60.98	0.84	95.98	
13105	64.62	26.94	43.02	1.19	95.98	
10659	132.22	90.74	258.54	43.13	95.94	
106	59.79	20.18	35.91	1.62	95.89	
2697	1226.39	358.41	1822.86	80.35	95.85	
10217	310.63	90.28	201.72	7.38	95.85	
22658	275.00	67.48	237.14	2.73	95.85	
3417	445.05	113.12	533.98	7.67	95.81	
15446	369.04	93.58	428.95	2.94	95.81	
2847	62.77	31.28	81.96	1.81	95.81	
2469	1383.42	482.22	963.09	26.57	95.81	
17175	619.05	156.40	867.90	59.01	95.77	
4386	56.38	26.60	45.57	0.72	95.77	
21491	110.61	27.92	90.43	1.09	95.77	
20844	843.25	294.01	1309.53	64.24	95.72	
9370	773.72	201.87	676.25	8.78	95.72	
15130	274.69	83.28	262.79	4.56	95.64	

Timepoint(PAN s): 163 lus			Dog. K	921-5039WO o. 1793397.1
ldendifter –	NonToxMean	NonToxeD:	ToxMean ::	TOXSD	LDAScore .
6100	-24.28	62.27	70.39	19.04	99.79
2048	31.63	20.68	119.91	5.57	99.79
24024	32.82	21.87	144.74	15.76	99.79
14722	658.73	219.20	1717.84	115.16	99.74
21651	31.52	24.04	198.29	19.08	99.74
15401	75.65	26.38	220.16	14.55	99.70
15503	124.16	37.19	305.20	27.55	99.70
7090	16.70	14.04	114.99	16.69	99.70
10893	-30.66	62.67	46.11	19.29	99.70
23780	24.98	35.81	102.35	16.07	99.66
16726	944.95	205.53	460.40	16.03	99.61
14430	33.41	27.29	289.53	65.58	99.61
15434	132.29	53.11	374.76	31.01	99.53
17198	812.71	267.14	249.19	36.68	99.53
15437	38.33	22.07	168.45	19.74	99.53
225	123.21	34.21	276.57	21.70	99.49
3584	64.34	31.89	290.58	28.71	99.49
9084	46.92	17.70	134.77	13.50	99.49
20736	331.77	100.84	748.89	43.21	99.49
19374	-8.73	12.21	63.84	14.49	99.44
20699	94.34	231.80	255.20	18.42	99.44
22816	23.08	15.16	83.31	4.90	99.44
23297	384.07	96.58	840.46	53.16	99.44
15002	140.16	98.17	338.90	19.21	99.44
3269	75.61	36.41	365.17	72.48	99.44
23773	212.24	81.04	629.63	48.29	99.40
23992	5.27	6.59	56.95	14.22	99.36
17682	673.28	176.72	223.65	27.55	99.36
3079	34.49	46.53	164.24	20.61	99.36
23778	68.50	31.99	270.34	37.08	99.36
19006	642.56	208.65	1445.68	75.63	99.31
20887	793.69	313.68	218.40	48.36	99.31
6039	310.04	86.04	693.54	28.37	99.27
15003	35.50	95.84	254.52	23.59	99.27
17227	873.36	191.79	476.74	33.25	99.27
1801	97.51	29.11	207.98	12.46	99.27
10015	231.53	75.18	560.75	25.13	99.27
12683	60.85	40.62	505.60	124.63	99.23
16675	33.50	33.48	151.49	16.04	99.23
1809	36.19	128.09	267.84	32.82	99.23
15981	75.09	25.95	290.62	83.27	99.23
2637	124.84	42.73 37.28	270.38 252.45	22.27 25.44	99.23 99.23
2284	99.14				99.23
11338 16425	39.20	31.91 26.43	342.99 256.34	92.86 69.36	99.19
9212	19.78 1017.57	301.78	2309.86	189.18	99.19
13977	281.63	142.69	11144.78	134.21	99.19
3572	33.77	17.83	169.37	47.47	99.19
22079	78.37	356.23	216.47	49.64	99.19
13974	298.69	149.15	842.98	65.09	99.19
24564	568.47	175.61	176.63	24.26	99.19
18603	-45.88	67.13	192.24	18.99	99.14
2010	33.66	315.34	180.93	36.19	99.14
17501	31.72	17.98	94.02	6.17	99.14
23376	14.35	15.90	95.17	30.08	99.14

Time point	PAN s): 168 hrs		Aity: Docket No. 44921-5039W0 Doc. No. 1793397.1			
ldentifier"	NonforMean	NontoxSD	ToxMean :	ToxSD	LD/AScore	
6517	231.39	155.95	730.29	101.68	99.14	
20599	22.90	12.70	74.26	5.67	99.14	
23377	61.42	52.07	359.41	62.64	99.14	
14405	459.81	287.63	2264.76	369.05	99.14	
1564	38.78	169.92	459.31	64.11	99.10	
16676	38.17	26.57	134.01	11.56	99.10	
21695	59.83	35.43	188.55	18.05	99.10	
20886	719.56	292.49	116.56	44.10	99.10	
17151	16.05	62.27	445.34	138.38	99.06	
770	798.22	219.65	310.69	35.90	99.06	
7262	1116.47	390.83	2517.22	160.89	99.06	
23981	809.28	155.43	1417.26	153.29	99.06	
22552	314.65	94.11	570.75	29.08	99.06	
20709	162.15	69.62	483.04	60.17	99.06	
7585	-91.46	37.80	142.97	84.57	99.01	
15853	5.82	59.90	258.64	131.89	99.01	
22592	233.64	160.86	1028.66	111.76	98.97	
4445	514.32	117.31	988.03	63.78	98.97	
4086	14.01	16.98	152.91	75.32	98.97	
21509	88.80	92.91	448.20	56.16	98.97	
24651	100.64	23.29	204.63	19.93	98.97	
7101	389.56	671.41	1218.47	165.33	98.93	
15851	200.60	148.70	784.83	83.88	98.93	
23769	-6.25	8.68	29.51	10.78	98.93	
15504	128.87	50.45	417.46	77.64	98.93	
1613	-7.12	28.04	117.88	29.33	98.93	
15438	65.66	32.54	171.12	14.45	98.93	
1460	198.26	83.19	563.02	70.47	98.93	
13976	449.17	351.62	2551.95	498.75	98.93	
20903	69.41	42.80	397.90	116.45	98.89	
23123	310.66	118.21	861.54	70.50	98.89	
15790	45.93	25.16	134.46	10.51	98.89	
21391	213.21	127.55	742.74	58.03	98.89	
455	137.60	153.63	269.32	29.16	98.89	
20772	149.52	40.53	313.83	28.96	98.89	
19275	726.18	191.87	1580.61	140.07	98.84	
17149	81.87	40.60	328.14	60.26	98.84	
19243	76.71	48.71	359.48	105.52	98.84	
17197	1903.25	828.79	668.38	83.11	98.84	
11891	-15.63	11.78	25.77	9.49	98.84	
4640	80.30	30.25	184.62	15.77	98.80	
4569	0.89	6.39	34.18	10.37	98.80	
2629	25.55	23.81	70.08	4.98	98.80	
18529	202.36	80.95	572.14	48.95	98.80	

TABLE 5Y: PAN Ally, Docket No. 44921-5039					
	(s): 6, 24 hrs	No-53	F-~00		9. 1793397.1
410	1099.85	252.82	487.89	71.66	99.01
1137	58.38	18.64	128.72	40.53	97.55
18322	2669.23	812.57	1422.54	109.34	97.47
15433	67.21	32.38	147.81	13.69	97.04
8990	278.32	77.85	504.55	67.10	96.35
23115	565.35	169.43	1013.53	113.02	96.22
1460	199.21	86.34	300.32	24.45	95.97
335	96.10	45.37	194.53	15.33	95.92
2866	637.05	212.44	1020.97	43.03	95.75
15701	37.69	16.68	82.72	11.76	95.75
16853	67.54	23.29	107.06	8.27	95.58
17693	1257.20	377.49	705.99	42.86	95.41
6250	490.36	103.66	731.52	54.23	95.36
19327	89.22	29.33	144.42	10.71	95.28
21977	88.79	52.26	163.28	17.00	95.11
1962	33.32	26.88	74.11	7.81	95.06
19080	75.82	57.08	219.58	56.51	95.06
13598	349.80	109.75	578.05	60.94	95.02
11524	-14.25	24.65	43.36	19.72	95.02
729	90.40	35.70	171.30	19.74	94.98
15552	175.81	40.06	237.25	9.67	94.98
4312	77.28	38.83	150.22	19.98	94.94
18996	137.86	44.69	250.58	35.37	94.85
17411	79.24	54.48	184.11	25.15	94.76
20752	13.07	11.34	26.51	2.40	94.76
11445	436.52	103.96	686.40	101.03	94.42
17755	393.23	102.51	214.27	40.93	94.42
19077	208.94	51.22	306.89	24.65	94.25
21355	375.05	118.77	619.86	88.05	94.21
6454	239.78	78.25	421.17	60.75	94.16
21092	392.44	225.02	575.95	55.98	94.12
11618	414.89	130.79	694.32	108.18	94.03
18338	72.29	20.32	112.46	9.87	94.03
15050	637.68	181.55	464.30	24.18	94.03
1608	12.70	29.99	54.74	9.39	93.99
24539	613.90	163.40	344.65	53.69	93.99
5900	265.48	65.87	137.54	40.46	93.99
13239	109.63	48.60	209.05	26.39	93.99
5163	20.12	13.08	44.33	7.38	93.82
21130	78.12	26.32	110.79	5.99	93.73
2236	140.10	38.70	215.23	16.78	93.69
5967	1225.74	339.49	1876.92	202.57	93.65
9799	145.09	50.35	241.50	35.40	93.56
16205	940.52	264.48	668.73	24.99	93.56
456	869.99	317.52	515.38	44.20	93.56
25097	4.68	10.99	25.79	8.52	93.52
12020	167.47	60.50	285.05	31.13	93.52
5924	162.21	46.91	256.59	29.67	93.52
4716	156.93	40.11	238.10	25.26	93.48
8339	432.55	130.90	704.91	90.15	93.43
5561	167.91	55.87	283.83	41.10	93.39
15112	1549.68	531.98	736.99	1111.89	93.39
		1	1077.40	67.94	93.35
7278	1467.27	341.78 29.56	1116.16	20.84	93.30
7108	53.62	38.21	153.60	24.20	93.30
4956	79.01		1		93.26
20404	53.75	46.18	126.25	22.59	J93.20

TABLE 5Y	's PAN		Ality. Dod	ket No. 449	021-50:9W 9. 179:397.1
	(S): 6,24 hrs NonToxMean	l Management	lstM		
					93.22
6049	621.00	123.14	541.13	9.05	93.22
9109	1021.28	337.87	723.36	26.27	93.22
4393	1268.21	369.74	824.30	42.57	93.13
18541	1081.46	298.74	758.66	24.93	
15556	208.38	83.99	333.56	45.17	93.09
4589	1213.38	369.83	790.80	42.48	93.09
22271	273.68	65.98	411.42	75.15	93.05
21423	1355.28	427.17	786.98	56.19	93.05
6506	234.14	60.71	335.18	21.90	93.05
4290	94.34	28.53	147.53	19.36	93.05
20350	155.10	50.13	248.15	31.72	93.05
714	46.67	31.34	115.67	27.05	93.05
1485	93.58	57.35	198.96	32.55	93.00
18433	19.39	46.95	105.10	24.21	93.00
17567	1185.38	364.09	750.70	33.02	93.00
24615	817.72	230.83	583.06	24.71	92.96
15742	33.17	14.60	56.07	3.60	92.96
20766	56.29	20.07	93.49	10.52	92.92
15209	164.94	33.20	220.88	12.45	92.92
20879	87.79	50.58	180.08	38.06	92.88
19408	2011.27	675.11	1014.96	97.93	92.88
1247	1309.18	501.52	569.36	138.91	92.83
15299	88.50	61.88	119.48	12.69	92.83
11377	116.76	30.96	177.28	17.37	92.79
9037	22.39	16.20	56.28	11.77	92.79
24390	162.85	113.82	331.70	49.97	92.79
1550	30.09	69.61	60.64	8.26	92.79
1300	156.68	44.76	234.60	19.19	92.75
1585	67.71	33.38	128.60	16.01	92.70
25599	56.14	20.41	91.99	8.79	92.66
17524	1173.79	285.74	801.36	115.86	92.62
15122	420.86	89.65	588.05	45.81	92.58
25369	18.36	10.68	38.74	6.19	92.58
3886	58.10	28.84	117.74	22.59	92.53
643	61.37	32.91	119.52	18.91	92.53
14003	815.86	212.90	497.74	73.28	92.49
16164	1083.28	283.92	680.99	54.84	92.49
20864	1587.98	660.11	798.40	87.84	92.45
17742	1066.26	308.06	723.12	37.02	92.45
23248	37.67	17.23	53.27	3.03	92.45
17204	1732.06	656.42	888.59	86.53	92.45
24501	525.70	117.48	664.66	34.25	92.40
14125	128.41	50.32	217.38	25.88	92.36
			1474.27	231.26	92.36
5968	962.02	263.60	14/4.2/	231.20	92.30

TABLE 572: Timepoint(PAN <u>s)): 6, 24, 163</u> h	The state of the s	Aiiy. Doc		921-5039V. 0. 1793397.1
relitionebl	NonToxMean	NonToxSD	ToxiMean	ToxSD 🔻	LDAScore -
15433	66.69	31.46	164.15	24.65	97.20
1962	33.08	26.68	80.19	11.69	95.26
16122	116.83	40.09	224.79	57.88	94.96
1247	1312.82	499.50	526.34	127.30	94.87
16853	67.23	22.82	120.51	22.86	94.83
17693	1260.09	375.71	656.50	74.07	94.78
13239	108.92	47.30	236.00	58.32	94.57
15112	1553.39	530.07	717.14	110.65	94.35
18433	18.80	46.13	126.18	39.27	94.35
16121	109.21	58.36	264.16	84.47	94.18
15701	37.56	16.60	76.48	12.40	94.05
19077	208.57	51.01	301.25	21.97	94.01
4723	1721.81	624.78	922.60	68.61	93.58
25097	4.60	10.95	24.13	7.34	93.23
7278	1469.82	340.27	993.59	122.75	93.23
1159	888.08	229.69	532.18	82.33	93.19
15184	-80.88	50.16	76.70	82.34	93.07
456	871.69	317.12	499.51	56.17	92.97
1608	12.56	29.97	50.60	9.98	92.93
15437	38.06	21.84	118.82	46.29	92.69
225	122.71	33.55	235.76	49.22	92.51
14997	2605.00	1019.08	1218.22	195.95	92.46
15401	75.23	25.84	175.78	44.88	92.39
19408	2015.21	673.87	1049.53	100.77	92.20
18076	2552.08	1185.85	1116.73	165.36	91.90
18077	2630.74	1183.04	1067.76	130.98	91.72
15434	131.68	52.58	292.95	80.76	91.61
16211	2268.34	1131.65	761.93	145.25	91.51
11208	322.94	110.75	642.67	141.37	91.48
24651	100.37	23.09	169.86	33.73	91.48
16012	72.52	31.21	141.73	32.89	91.35
1542	928.57	263.76	548.27	84.14	91.34
14722	656.51	217.90	1314.74	367.97	91.31
4589	1215.64	368.99	748.52	72.66	91.21
17886	1506.14	491.91	843.64	96.88	91.16
17829	2232.91	1111.68	900.83	138.88	91.16
15673	804.12	164.51	1076.89	78.92	91.16
4312	76.92	38.49	154.79	22.22	91.09
2079	303.47	81.98	400.13	28.12	90.99
18694	53.10	48.50	177.05	53.91	90.96
18322	2673.81	811.21	1499.15	181.85	90.79
16168	350.70	206.64	479.34	84.79	90.78
17779	1958.43	783.05	911.49	158.95	90.73
8990	277.74	77.38	465.89	83.79	90.57
3434	322.35	138.05	614.67	136.42	90.57
18729	19.67	29.33	96.11	37.87	90.57
14003	817.64	211.58	458.34	85.08	90.53
22816	22.96	15.10	59.89	23.04	90.49
9799	144.50	49.47	258.44	54.08	90.49
1521	20.40	50.29	136.10	71.19	90.49
15886	302.74	61.73	436.98	44.28	90.45
16155	1084.95	423.63	632.04	54.95	90.43
1485	92.98	56.73	213.05	31.14	90.36
17204	1735.18	656.07	938.01	109.72	90.30
1203	4.95	18.70	44.90	22.64	90.27

TABLE 572: PAN			Atty, Docket No. 44921-5089W0			
	Timepolini(s): 6, 24, 163 livs				40. 1793397.1	
relitioned)	NonToxMean	NonToxSD	ToxMean	ToxSD		
19080	75.38	56.65	202.18	64.18	90.27	
1622	2078.05	898.00	838.34	144.28	90.26	
17712	858.75	187.72	604.46	57.40	90.26	
19407	1533.63	471.44	919.83	109.37	90.26	
4280	1275.34	435.07	698.85	97.31	90.22	
635	1254.34	431.22	723.36	97.99	90.17	
22554	544.56	150.27	343.47	69.26	90.17	
11524	-14.44	24.49	37.71	21.53	90.14	
21703	16.32	12.14	42.83	13.13	90.10	
14125	127.72	48.99	246.62	72.21	90.06	
17480	119.30	33.27	190.90	31.79	90.01	
23322	1169.31	283.61	760.72	114.86	89.96	
8234	49.37	21.69	92.92	31.52	89.93	
13682	176.92	61.79	309.18	50.37	89.89	
11321	175.44	50.38	286.20	55.16	89.89	
25253	291.91	64.62	414.09	38.27	89.71	
23852	262.19	102.61	480.31	78.92	89.71	
20781	57.13	21.07	88.06	13.32	89.70	
1582	16.67	13.15	45.26	13.10	89.67	
24696	68.64	47.19	170.78	42.12	89.67	
818	4144.91	2804.50	980.91	271.23	89.66	
9109	1022.78	337.79	702.34	58.34	89.66	
21708	36.51	16.01	73.22	15.22	89.63	
17549	1309.92	366.25	854.17	88.86	89.61	
16591	151.86	47.16	253.45	36.23	89.54	
17154	198.88	57.70	280.48	33.03	89.53	
14694	2332.83	1063.34	1038.43	256.02	89.48	
23109	2093.27	924.06	1002.79	195.59	89.48	
2236	139.93	38.69	200.68	21.94	89.48	
22661	1310.36	418.55	756.78	88.01	89.48	
7857	51.98	45.43	149.29	46.95	89.45	
20876	1681.76	611.63	958.77	99.56	89.44	
25468	2183.54	1132.38	823.42	189.02	89.44	
23215	106.03	29.48	169.54	21.53	89.41	
5900	265.96	65.56	144.68	40.70	89.41	
1811	10.93	26.49	65.20	23.39	89.37	
23709	2507.78	1197.86	1088.40	231.11	89.35	
495	135.24	83.30	303.90	57.27	89.32	
20998	161.29	73.65	276.37	48.32	89.32	
17194	46.75	27.72	107.07	19.66	89.32	
19327	88.98	29.10	144.83	20.18	89.28	
18101	81.85	27.96	121.65	14.02	89.27	
819	3112.24	1661.98	1015.34	283.90	89.27	
3015	2325.38	988.40	1058.04	289.77	89.22	
21355	374.22	118.21	598.37	91.02	89.20	

		-43	3-		
	AA: SEMUSTIN M(S): 163 hrs	E	Ally. Do		4921-5039\WO No. 1793397.1
negitien Joentifie		[Real molt]	∏oxMean		LDAScore
	-3.74	14.74	24.13	2.44	98.46
23830 2655	43.68	49.43	103.83	13.87	98.07
2055 22547	15.31	164.20	55.98	7.70	97.99
21893	51.15	34.39	119.77	15.62	97.86
7806	52.07	19.62	68.71	1.17	97.73
7785	1830.12	494.30	1054.17	67.70	97.69
25705	455.94	115.83	667.36	30.91	97.64
22050	3163.82	930.45	2122.73	41.59	97.47
21443	103.90	53.37	35.53	5.08	97.26
23981	812.67	160.21	626.66	12.41	97.13
5733	12.64	47.16	51.31	11.73	97.13
8494	276.36	53.17	213.99	3.26	97.13
	25.65	33.31	2.82	0.93	97.13
8901 7023		77.19	291.07	3.81	97.09
	364.88 113.25	70.77	275.16	28.92	97.04
18353	1129.65		11127.48	12.54	97.04
3027		333.83 1495.61	2320.07	89.57	96.92
8177	3894.73			72.18	96.83
19398	4332.61	2167.49	2566.57 159.26	19.40	96.83
23626	71.43	37.20		13.70	96.74
18650	804.69	246.36	518.86 3041.20	196.69	96.62
15627	5054.50	1946.06			
20757	413.88	222.48	607.95	51.66	96.53
2410	11.25	10.57	25.63	2.99	96.49
7274	576.90	150.70	364.54	16.30	96.44
26147	1030.83	225.21	645.26	49.87	96.32
3359	119.66	57.61	290.85	65.10	96.27
11994	104.13	23.52	63.18	4.39	96.06
7867	64.06	34.54	163.76	30.00	96.02
3701	90.19	44.20	111.89	2.06	95.93
23800	32.07	18.29	77.92	15.10	95.84
1684	2810.78	1628.26	2548.66	84.69	95.84
19162	2230.57	613.60	1498.90	56.20	95.84
14958	138.49	48.09	181.53	4.13	95.80
11454	238.93	79.65	379.87	38.25	95.76
7764	6.68	15.73	27.19	3.33	95.76
19993	2327.93	570.40	1514.68	94.53	95.76
9407	350.93	216.90	167.34	16.19	95.76
15600	772.34	211.63	480.10	25.98	95.76
22662	137.98	35.08	116.24	1.67	95.72
18142	1994.46	839.22	1965.70	25.18	95.63
7278	1466.77	341.68	1037.78	55.91	95.59
1685	9177.09	7104.79	4192.02	442.45	95.59
15136	720.68	229.89	618.26	7.44	95.50
17248	2563.75	537.86	1741.86	94.90	95.46
15301	38.66	68.20	52.95	4.33	95.46
24627	5024.80	1714.05	3187.73	133.28	95.46
16678	192.60	90.16	295.43	9.43	95.37
21798	2088.65	441.72	1574.36	40.27	95.37
3582	396.47	105.41	299.17	10.26	95.29
22618	28.62	11.98	54.72	9.80	95.24
20295	23.12	13.39	-1.42	5.25	95.20
10269	1868.08	422.77	1408.13	42.28	95.16
7975	744.31	175.27	709.93	7.00	95.12
3090	308.90	80.19	389.68	9.44	95.07
8163	105.92	55.18	246.10	47.62	95.03

TABLE SA	A: SEMUSTINE (S): 168 hrs	tri er enrigeres	Ally, Do	cket No. 44	1921-5089///0
aniomenalii	(s)): 168 hrs		73.4	Doc. 1	lo. 179839 7.1
ldendfler	NonToxMean :	NonToxSD	ToxMean	ToxSD	LDAScore
7337	130.17	35.83	72.98	11.19	95.03
5689	12.93	20.23	44.79	8.55	94.99
22737	204.61	104.35	375.42	54.48	94.94
4640	80.47	30.73	144.33	17.03	94.82
10819	1187.21	363.05	1125.76	18.01	94.82
3269	76.54	40.97	147.76	18.49	94.77
18468	90.58	33.19	99.23	1.85	94.77
6204	39.16	12.29	53.01	1.67	94.77
19372	61.97	54.00	9.74	2.20	94.73
21628	116.51	45.00	204.96	26.35	94.69
3075	523.79	180.19	291.98	21.30	94.69
8015	52.45	36.45	80.79	2.30	94.64
20105	80.00	53.71	27.37	3.94	94.64
18830	5928.33	2277.75	3404.34	229.07	94.60
12639	1231.40	359.71	1328.85	25.82	94.60
2697	1227.29	359.86	1375.72	25.94	94.56
2326	1442.32	399.30	866.19	98.10	94.56
9180	82.53	35.16	153.13	21.13	94.47
22838	1383.85	399.86	912.16	54.21	94.47
23509	2234.19	462.51	1771.07	42.02	94.47
23709	2494.50	1202.87	2200.91	59.43	94.34
2912	2618.16	709.02	1838.26	100.99	94.34
20944	1191.93	395.05	1322.96	34.77	94.30
25589	154.92	38.08	184.54	2.96	94.30
18800	2730.47	1166.13	1296.14	149.09	94.26
2311	22.96	16.01	54.56	12.04	94.17
410	1097.48	256.27	795.24	44.26	94.17
17885	246.73	58.07	148.75	13.01	94.13
21458	229.64	104.63	360.82	32.46	94.13
22069	689.12	145.53	514.75	17.18	94.13
3584	65.13	35.08	106.58	9.75	94.13
5968	966.61	265.81	609.81	49.16	94.04
19525	27.97	21.00	56.08	10.20	94.04
18757	250.38	72.47	327.68	9.19	94.00
19768	711.98	179.77	874.96	22.13	94.00
7392	29.06	15.76	62.61	15.61	94.00
10267	2119.14	891.88	2056.03	48.77	94.00
23190	70.87	28.52	103.79	5.80	93.96
25676	218.79	80.22	97.09	19.62	93.96

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nmebon	nt(s): 24 hrs 7 Nonvoxivean	10.1/23231/2323-401		Dioice	. No. 1798897.1
ldentitiei				11.81	199.83
13158	464.71 31.95	105.53 313.46	185.18 578.42	46.88	99.79
2010			792.83	41.28	99.74
15535	445.86	80.88 147.30	1395.56	106.27	99.70
1564	34.77		328.14	43.97	99.70
4452	989.99	248.64 50.39	113.73	16.34	99.61
4450	282.32		1604.45	209.39	99.61
20700	81.27	404.57 44.26	81.56	7.25	99.57
20481	208.68 666.44	155.98	1348.56	47.39	99.53
25686	1559.28	402.43	636.16	43.80	99.53
5695	13.20	160.91	187.40	49.13	99.53
6477	352.23	77.89	621.28	16.27	99.49
17101 15175	318.83	49.56	153.62	12.70	99.49
20699	90.33	220.48	1192.64	109.71	99.49
	22.55	383.52	251.88	88.78	99.44
6478		45.62	14.15	2.76	99.44
725	117.71	54.99	17.82	6.85	99.44
645	151.16		465.13	55.22	99.44
3987	192.50	47.15	319.39	29.30	99.40
21947	752.67	136.25	1		99.40
17291	1389.57	412.30	482.86	101.58	99.40
238	227.68	51.22	496.72	69.45	
20887	794.18	312.69	103.79	42.68	99.36
1558	203.57	58.77	553.28	64.74	99.36
17661	255.29	58.73	540.05	70.79	99.36
16947	320.42	79.17	77.47	19.86	99.36
16204	752.56	177.92	1702.82	145.45	99.36
18906	255.45	83.89	55.16	8.36	99.36
17357	269.77	81.23	2.53	12.73	99.31
17514	859.90	174.50	359.55	37.03	
16416	22.83	18.51	84.23	3.30	99.27
2085	688.34	191.75	339.59	20.82	99.27
7101	380.51	644.25	3330.36	683.13	99.27
18278	1896.72	507.81	639.93	108.91	99.27
25718	392.38	80.21	851.19	103.08	99.23
1478	352.13	79.44	88.92	22.22	99.23
9621	292.01	59.49	563.88	51.66	99.23
23596	1491.59	375.55	487.23	90.69	99.23
20884	637.92	258.49	19.93	9.59	99.23
10016	207.78	73.01	590.63	39.54	99.23
7665	282.42	91.44	861.31	113.78	99.19
472	661.53	177.31	345.51	23.33	99.19
18597	521.43	162.83	914.02	31.92	99.19
4222	545.41	102.20	1014.25	79.67	99.19
20886	719.84	291.88	50.84	41.81	99.19
1884	178.24	37.12	331.54	22.95	99.19
1422	297.77	88.40	75.68	10.49	99.19
17104	449.21	107.90	1060.31	149.95	99.14
17907	1319.66	300.38	623.40	49.85	99.14
17284	229.07	60.08	77.02	16.01	99.14
8981	142.60	87.32	716.17	136.32	99.14
4451	290.54	64.21	76.77	27.51	99.14
15955	751.14	196.31	146.28	35.67	99.10
16419	306.72	58.43	562.97	56.12	99.10
16627	97.46	34.41	213.43	14.92	99.10
22592	234.18	163.38	903.10	59.07	99.10
19824	224.78	67.21	32.81	20.42	99.10

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Timenoin	((s): 24 hrs			Doc.	No. 1793397.1	
leenuiter		INOUI OXX		ା ପ୍ରକ୍ଷର 🏥	TEDY/2000	
9905	674.04	138.26	279.49	36.23	99.10	
15002	139.13	94.64	580.42	72.97	99.10	
812	157.04	34.99	39.35	14.07	99.10	
15193	172.71	60.15	463.80	40.10	99.10	
11205	596.04	192.04	155.98	25.86	99.10	
17535	246.41	64.45	529.14	76.63	99.06	
4360	330.80	65.83	138.60	23.78	99.06	
5667	742.84	172.36	1513.63	136.63	99.06	
20056	309.09	59.92	81.20	26.34	99.06	
11081	352.98	113.10	740.48	49.43	99.06	
3916	738.87	186.54	212.10	43.88	99.06	
1801	97.49	29.02	212.29	21.51	99.06	
25317	15.28	68.43	72.14	39.26	99.06	
24501	524.69	114.42	955.66	63.01	99.06	
19508	100.97	41.09	11.48	5.01	99.06	
19769	63.82	80.07	487.54	129.83	99.01	
25546	473.54	139.58	193.05	34.65	99.01	
10878	951.05	250.12	1714.95	105.09	99.01	
10819	1183.30	358.76	2036.62	53.76	99.01	
18507	701.19	159.97	1396.26	108.52	98.97	
2915	87.36	36.13	206.79	19.36	98.97	
15003	34.41	91.73	507.05	83.25	98.97	
2583	369.08	148.55	848.08	90.80	98.97	
9620	529.25	111.98	1071.29	125.35	98.97	
19298	364.57	96.71	819.01	116.31	98.97	
6614	478.70	150.19	97.33	35.20	98.97	
15986	320.13	83.84	46.97	24.78	98.97	
13646	727.39	191.95	1634.83	174.87	98.93	
240	184.44	53.70	394.90	58.94	98.93	
16675	33.18	31.66	226.87	74.32	98.93	
11968	297.46	82.47	94.28	17.16	98.93	
17540	561.49	199.04	1593.99	230.39	98.93	
19161	1068.85	309.26	2020.78	120.96	98.93	
23924	208.78	62.50	465.78	53.25	98.93	
4482	227.17	73.35	67.19	17.67	98.93	
5199	611.50	182.04	158.54	39.19	98.93	
117	21.06	17.59	-12.77	2.61	98.89	
22536	1715.17	473.49	3319.42	278.50	98.89	
17100	819.32	235.74	1519.10	85.28	98.89	
14332	537.23	133.39	1143.62	152.21	98.89	
5232	365.72	93.58	160.50	9.58	98.89	
10561	78.49	36.71	188.47	14.45	98.89	
15098	185.41	66.49	32.78	6.96	98.89	
3401	14.24	21.17	144.41	52.94	98.89	

	: SULFADIAZIN 3): 3, 6 hrs		921-5039WO o. 1793397.1		
Maniffar Identiffar	MonToxMean	NonToxSD'			
2459	327.82	133.31	1186.99	180.16	99.44
985	18.91	23.49	435.93	226.01	99.36
6054	24.46	35.46	463.67	144.11	99.18
21546	-60.81	78.07	461.08	143.78	99.14
11259	101.27	105.67	1117.26	304.40	99.05
223	10.81	16.39	121.57	31.76	99.05
13745	27.17	26.30	236.39	104.08	99.05
8304	263.66	68.46	104.04	18.52	99.05
2154	54.19	102.13	766.15	271.93	98.93
6585	635.65	314.22	2778.88	622.70	98.80
17506	33.02	28.55	170.00	54.28	98.80
2153	179.20	124.41	1217.91	474.19	98.75
23872	47.29	87.73	387.10	112.60	98.71
15301	35.41	55.60	466.69	146.16	98.67
15192	164.68	117.96	858.98	208.00	98.67
17327	209.40	86.17	531.34	65.50	98.62
15300	131.96	108.91	967.46	264.07	98.62
7700	80.97	38.43	377.20	103.65	98.58
13930	128.33	69.33	520.74	123.87	98.54
10659	128.90	79.62	602.71	192.55	98.54
7197	195.08	78.35	557.25	113.13	98.37
19060	150.81	79.81	399.19	134.24	98.32
15299	86.11	52.88	422.06	161.10	98.28
355	10.95	13.39	102.92	40.18	98.28
7196	178.85	80.11	517.52	81.96	98.28
25730	196.00	64.30	513.70	106.95	98.24
5356	-12.88	18.58	46.77	19.05	98.19
19657	3.19	17.88	84.23	34.61	98.15
25084	-12.38	22.65	46.65	25.44	98.11
1501	58.67	53.51	250.18	120.56	98.02
17908	62.13	58.89	240.27	66.39	97.98
16314	43.27	38.63	240.75	61.16	97.94
2555	95.85	44.10	292.92	71.25	97.94
21682	-35.82	31.41	50.22	26.54	97.89
5384	40.19	41.73	218.67	41.90	97.89
22626	83.22	71.71	465.98	192.17	97.85
11483	64.30	44.10	197.71	47.03	97.85
16053	229.00	93.51	658.28	200.15	97.81
4045	93.63	47.19	244.15	46.42	97.81
804	454.28	182.83	934.61	83.33	97.76
8874	118.37	60.25	305.67	73.98	97.76
23314	62.72	247.80	1403.77	543.62	97.76
21	69.73	31.90	161.45	47.41	97.76
22681	218.11	182.12	981.97	310.38	97.72
7471	219.05	63.97	421.52	40.74	97.72
13240	475.09	362.91	2390.71	790.75	97.68
1340	192.32	49.81	108.32	15.13	97.64
16312	54.74	33.76	197.76	42.24	97.59
11876	107.52	40.42	242.85	40.07	97.46
20161	38.17	33.28	150.28	32.80	97.46
15996	49.41	38.61	170.53	42.38	97.42
12978	105.07	45.80	334.16	95.16	97.42
14179	49.88	26.28	132.84	24.19	97.38
12534	87.91	36.00	223.28	65.14	97.38
21779	147.51	35.16	262.54	35.74	97.29

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TABLE SCC: SULFADIAZINE ATTY. Docket No. 44921-5039WC							
ldendifier	NonToxMean:	NonToxSD	ToxWean	ToxSD /	LDAScore		
11979	210.67	69.55	85.98	17.52	97.25		
17161	1152.09	406.69	2288.34	334.10	97.25		
11434	326.14	103.27	663.21	132.01	97.12		
6844	124.34	58.18	17.91	11.06	97.12		
22	53.08	46.64	206.89	59.81	97.12		
3464	165.60	53.31	329.80	63.59	97.08		
3434	322.01	136.13	755.72	108.22	97.08		
22248	243.91	118.15	625.27	179.46	97.03		
9423	937.34	251.31	1765.27	274.26	97.03		
21683	34.05	23.48	105.62	26.60	96.95		
16527	51.16	16.84	24.15	3.25	96.90		
4512	1.04	32.53	73.93	19.11	96.86		
4725	81.40	55.94	245.03	66.25	96.78		
21239	114.57	59.35	256.57	37.93	96.78		
17329	213.88	104.30	531.44	105.41	96.65		
13467	37.67	24.44	111.98	36.67	96.56		
15089	179.58	77.35	389.68	58.33	96.56		
3454	86.52	35.67	198.35	31.79	96.47		
20523	547.69	149.96	965.84	112.80	96.43		
15058	197.84	70.39	377.22	46.94	96.35		
14492	613.89	164.78	1076.44	162.75	96.35		
12999	128.09	33.18	203.14	17.45	96.30		
22967	163.84	54.49	255.51	13.19	96.26		
18008	1.51	16.75	42.22	12.38	96.26		
24640	177.45	57.95	357.46	73.86	96.26		
7913	55.20	20.69	108.16	18.70	96.17		
2625	141.92	31.66	87.09	9.03	96.17		
3049	201.89	100.36	461.30	105.24	96.17		
18011	31.95	28.60	105.17	25.57	96.13		
23834	71.40	23.40	137.38	25.69	96.09		
19818	32.37	21.21	85.97	15.20	96.04		
12673	37.51	22.58	99.74	25.41	96.04		
15377	25.91	15.53	56.38	6.78	96.00		
12745	330.67	112.37	623.69	81.74	96.00		
14111	150.33	66.27	327.97	47.79	96.00		
11157	681.75	178.91	356.49	80.44	96.00		
23166	134.04	62.51	266.66	54.48	95.96		
3519	375.54	125.21	690.15	176.36	95.87		
16124	420.65	131.40	193.12	32.21	95.83		
17800	204.49	42.25	108.97	22.01	95.79		
8639	368.08	97.97	620.44	125.57	95.79		
25090	83.00	43.32	203.07	48.58	95.79		
15191	2005.55	1208.87	3819.43	343.39	95.74		
7469	286.95	67.51	149.39	36.61	95.70		
21238	-17.22	39.01	55.57	20.45	95.70		

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WE CLAIM:

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- 1. A method of predicting at least one toxic effect of a compound, comprising:
- (a) preparing a gene expression profile of a tissue or cell sample exposed to the compound; and
- 5 (b) comparing the gene expression profile to a database comprising at least part of the data or information of Tables 1-5.
 - 2. A method of claim 1, wherein the gene expression profile prepared from the tissue or cell sample comprises the level of expression for at least one gene.
 - 3. A method of claim 2, wherein the level of expression is compared to a Tox Mean and/or NonTox Mean value in Tables 5-5CC.
- 4. A method of claim 3, wherein the level of expression is normalized prior to comparison.
 - 5. A method of claim 1, wherein the database comprises substantially all of the data or information in Tables 5-5CC.
- A method of predicting at least one toxic effect of a compound, comprising:
 - (a) detecting the level of expression in a tissue or cell sample exposed to the compound of two or more genes from Tables 1-5; wherein differential expression of the genes in Tables 1-5 is indicative of at least one toxic effect.
- 7. A method of predicting the progression of a toxic effect of a compound, comprising:
 - (a) detecting the level of expression in a tissue or cell sample exposed to the compound of two or more genes from Tables 1-5; wherein differential expression of the genes in Tables 1-5 is indicative of toxicity progression.
 - 8. A method of predicting the renal toxicity of a compound, comprising:
 - (a) detecting the level of expression in a tissue or cell sample exposed to the

compound of two or more genes from Tables 1-5; wherein differential expression of the genes in Tables 1-5 is indicative of renal toxicity.

- 9. A method of identifying an agent that modulates the onset or progression of a
 5 toxic response, comprising:
 - (a) exposing a cell to the agent and a known toxin; and
 - (b) detecting the expression level of two or more genes from Tables 1-5; wherein differential expression of the genes in Tables 1-5 is indicative of toxicity.
- 10 10. A method of predicting the cellular pathways that a compound modulates in a cell, comprising:

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- (a) detecting the level of expression in a tissue or cell sample exposed to the compound of two or more genes from Tables 1-5; wherein differential expression of the genes in Tables 1-5 is associated the modulation of at least one cellular pathway.
- 11. The method of any one of claims 6-10, wherein the expression levels of at least 3 genes are detected.
- 12. The method of any one of claims 6-10, wherein the expression levels of at least 4 genes are detected.
 - 13. The method of any one of claims 6-10, wherein the expression levels of at least 5 genes are detected.
- 25 14. The method of any one of claims 6-10, wherein the expression levels of at least 6 genes are detected.
 - 15. The method of any one of claims 6-10, wherein the expression levels of at least 7 genes are detected.
 - 16. The method of any one of claims 6-10, wherein the expression levels of at least 8 genes are detected.

- 17. The method of any one of claims 6-10, wherein the expression levels of at least 9 genes are detected.
- 5 18. The method of any one of claims 6-10, wherein the expression levels of at least 10 genes are detected.
 - 19. A method of claim 6 or 7, wherein the effect is selected from the group consisting of nephritis, kidney necrosis, glomerular and tubular injury, and focal segmental glomerulosclerosis.

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- 20. A method of claim 8, wherein the renal toxicity is associated with at least one kidney disease pathology selected from the group consisting of nephritis, kidney necrosis, glomerular and tubular injury, and focal segmental glomerulosclerosis.
- 21. A method of claim 10, wherein the cellular pathway is modulated by a toxin selected from the group consisting of cephaloridine, cisplatin, puromycin aminonucleoside (PAN), bromoethylamine hydrobromide (BEA), gentamicin, ifosfamide, cyclophosphamide, carboplatin, AY-25329, indomethacin, acyclovir, citrinin, mercuric chloride, diflunisal, cidofovir, pamidronate, lithium, hydralazine, colchicine, sulfadiazine, and adriamycin.
- 22. A set of at least two probes, wherein each of the probes comprises a sequence that specifically hybridizes to a gene in Tables 1-5.
- 23. A set of probes according to claim 22, wherein the set comprises probes that hybridize to at least 3 genes.
- 24. A set of probes according to claim 22, wherein the set comprises probes that 30 hybridize to at least 5 genes.
 - 25. A set of probes according to claim 22, wherein the set comprises probes that

hybridize to at least 7 genes.

26. A set of probes according to claim 22, wherein the set comprises probes that hybridize to at least 10 genes.

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- 27. A set of probes according to any one of claims 22-26, wherein the probes are attached to a solid support.
- 28. A set of probes according to claim 27, wherein the solid support is selected from the group consisting of a membrane, a glass support and a silicon support.
 - 29. A solid support comprising at least two probes, wherein each of the probes comprises a sequence that specifically hybridizes to a gene in Tables 1-5.
- 15 30. A solid support of claim 29, wherein the solid support is an array comprising at least 10 different oligonucleotides in discrete locations per square centimeter.
 - 31. A solid support of claim 29, wherein the array comprises at least about 100 different oligonucleotides in discrete locations per square centimeter.

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- 32. A solid support of claim 29, wherein the array comprises at least about 1000 different oligonucleotides in discrete locations per square centimeter.
- A solid support of claim 29, wherein the array comprises at least about 10,000
 different oligonucleotides in discrete locations per square centimeter.
 - 34. A computer system comprising:
 - (a) a database containing information identifying the expression level in a tissue or cell sample exposed to a renal toxin of a set of genes comprising at least two genes in Tables 1-5; and
 - (b) a user interface to view the information.

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- 35. A computer system of claim 34, wherein the database further comprises sequence information for the genes.
- 36. A computer system of claim 34, wherein the database further comprises
 5 information identifying the expression level for the set of genes in the tissue or cell sample before exposure to a renal toxin.
 - 37. A computer system of claim 34, wherein the database further comprises information identifying the expression level of the set of genes in a tissue or cell sample exposed to at least a second renal toxin.
 - 38. A computer system of any of claims 34-37, further comprising records including descriptive information from an external database, which information correlates said genes to records in the external database.
 - 39. A computer system of claim 38, wherein the external database is GenBank.
 - 40. A method of using a computer system of any one of claims 34-37 to present information identifying the expression level in a tissue or cell of at least one gene in Tables 1-5, comprising:

comparing the expression level of at least one gene in Tables 1-5 in a tissue or cell exposed to a test agent to the level of expression of the gene in the database.

- 41. A method of claim 40, wherein the expression levels of at least two genes are compared.
 - 42. A method of claim 40, wherein the expression levels of at least five genes are compared.
- 30 43. A method of claim 40, wherein the expression levels of at least ten genes are compared.

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- 44. A method of claim 40, further comprising the step of displaying the level of expression of at least one gene in the tissue or cell sample compared to the expression level when exposed to a toxin.
- 5 45. A method of claim 9, wherein the known toxin is a renal toxin.

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- 46. A method of claim 42, wherein the renal toxin is selected from the group consisting of cephaloridine, cisplatin, puromycin aminonucleoside (PAN), bromoethylamine hydrobromide (BEA), gentamicin, ifosfamide, cyclophosphamide, carboplatin, AY-25329, indomethacin, acyclovir, citrinin, mercuric chloride, diflunisal, cidofovir, pamidronate, lithium, hydralazine, colchicine, sulfadiazine, and adriamycin.
 - 47. A method of any one of claims 6-10, wherein nearly all of the genes in Tables 1-5 are detected.
 - 48. A method of claim 47, wherein all of the genes in at least one of Tables 5-5CC are detected.
- 49. A kit comprising at least one solid support of any one of claims 29-33 packaged with gene expression information for said genes.
 - 50. A kit of claim 49, wherein the gene expression information comprises gene expression levels in a tissue or cell sample exposed to a renal toxin.
- 25 51. A kit of claim 50, wherein the gene expression information is in an electronic format.
 - 52. A method of any one of claims 6-10, wherein the compound exposure is *in vivo* or *in vitro*.
 - 53. A method of any one of claims 6-10, wherein the level of expression is detected by an amplification or hybridization assay.

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- 54. A method of claim 53, wherein the amplification assay is quantitative or semiquantitative PCR.
- 5 55. A method of claim 53, wherein the hybridization assay is selected from the group consisting of Northern blot, dot or slot blot, nuclease protection and microarray assays.
 - 56. A method of identifying an agent that modulates at least one activity of a protein encoded by a gene in Tables 1-5 comprising:
- 10 (a) exposing the protein to the agent; and

- (b) assaying at least one activity of said protein.
- 57. A method of claim 56, wherein the agent is exposed to a cell expressing the protein.
- 58. A method of claim 57, wherein the cell is exposed to a known toxin.
- 59. A method of claim 58 wherein the toxin modulates the expression of the protein.